Metabolic Dysfunction and Unabated Respiration Precede the Loss of Membrane Integrity during Dehydration of Germinating Radicles

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This study shows that dehydration induces imbalanced metabolism before loss of membrane integrity in desiccation-sensitive germinated radicles. Using a photoacoustic detection system, responses of CO2 emission and fermentation to drying were analyzed non-invasively in desiccation-tolerant and -intolerant radicles of cucumber (Cucumis sativa) and pea (Pisum sativum). Survival after non-invasively in desiccation-tolerant and -intolerant radicles of gravitate membrane damage induced by dehydration.

products resulting from imbalanced metabolism in seeds may ag-
transform infrared spectroscopy, acetaldehyde was found to disturb
appears to be associated with desiccation tolerance. Using Fourier

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result of the impairment of electron transport chains during
drying is thought to be the cause of peroxidative damage.
Biochemical studies have identified three factors contrib-
uting to a lethal accumulation of reactive O2 species in non-photosynthetic tissues (Nohl and Jordan, 1986; Cross and Jones, 1991; Skulachev, 1996): (a) redox states of the electron carriers inside mitochondria when ADP is limiting, (b) O2 availability, and (c) the absence of effective antioxidant systems.

Because desiccation-tolerant tissues do not exhibit symp-
toms of reactive O2 species-induced injury, they are en-
dowed with mechanisms controlling these three factors
during the loss of water. Although these factors are per-
ceived to be critical to achieve desiccation tolerance (Hen-
dry et al., 1992; Leprince et al., 1994, 1995; Leprince and Hoekstra, 1998), their relative importance and their control
during drying remain to be ascertained. In previous stud-
ies, we suggested that a coordinated down-regulation of metabolism in seeds early during drying may play an important role in avoiding oxidative stress conditions and/or accumulation of by-products of metabolism to toxic levels (Leprince and Hoekstra, 1998; Leprince et al., 1999). Down-regulation of metabolism appears to be an ancient and widespread regulatory mechanism that allows aerobes to withstand severe environmental stresses such as anoxia, freezing, and dehydration (Hand and Hardewig, 1996; Hardie et al., 1998). There is only limited evidence for metabolic depression in developing seeds (Kollöffel and Matthews, 1983) and somatic embryos (Tetteroo et al., 1995). The objective of this study was to assess whether desiccation tolerance of imbibed radicles of pea and cu-
cumber is correlated with a down-regulation of respiration.

In desiccation-tolerant seeds and pollen, several meta-
abolic parameters indicate that mitochondria experience a
decreasing O2 availability when tissues are dried below 0.6
g/g (g water/g dry weight) (Leprince and Hoekstra, 1998).
The cause for this desiccation-induced hypoxia is thought
to be an impeded diffusion of O2 as a result of the expo-
tential increase in viscosity during water loss (Leprince and Hoekstra, 1998). A characteristic feature of O2 defi-
ciency in plant metabolism is the shift from respiration to fermentation to maintain ATP levels and redox power,
resulting in the production of acetaldehyde and ethanol
(Kennedy et al., 1992; Drew, 1997). Anoxia and particularly
post-anoxic reoxygenation cause irreversible damage to
mitochondrial ultrastructure and energy metabolism (Drew, 1997; Zuckermann et al., 1997). A 1- to 2-h anoxic treatment given to 3-d-old germinated seeds of soybean induced a 50% loss of viability when returned to air (Van Toai and Bolles, 1991). The loss of viability of dry seeds during long-term storage has been correlated with a slow production of acetaldehyde (Zhang et al., 1995). An increased production of acetaldehyde and ethanol during seed imbibition is often a symptom of imbibitional injury or deterioration following storage, both leading to poor seed vigor or low germination rates (Woodstock and Taylorson, 1981a, 1981b; Gorecki et al., 1984). Aerobic fermentation occurs during the early hours of imbibition (Raymond et al., 1985; Kennedy et al., 1992). However, it is not known whether embryonic tissues produce acetaldehyde and ethanol during drying and whether there is a different pattern in the release of these volatiles in relation to desiccation tolerance. Therefore, this study compared the emission of ethanol and acetaldehyde during dehydration of imbibed seeds in relation to desiccation tolerance. Pea and cucumber seeds were chosen as representatives of starchy and oily seeds, respectively. In pea seeds, ATP production by alcoholic fermentation in normoxia may contribute up to 40% to the total energy production during early imbibition. In oily seeds, oxidative phosphorylation is the dominating ATP-synthesis pathway (Raymond et al., 1985). Furthermore, in germinated cucumber seeds that had lost their desiccation tolerance, this tolerance can conveniently be re-induced by a mild water stress for several days (Bruggink and van der Toorn, 1995).

A recurrent problem when one attempts to characterize metabolism in dehydrating seeds in relation to desiccation tolerance is the need for non-invasive and sensitive techniques that do not require water at some point in the analysis. This was overcome here by taking advantage of the sensitive photo-acoustic spectroscopy (PA) techniques available to study, on-line and non-invasively, the kinetics of emission of volatiles during dehydration. Because the PA setup is a flow-through system, the time response is fast, thereby avoiding the resort to head-space analysis as in conventional gas chromatography (GC) methods (Harren and Reuss, 1997; Zuckermann et al., 1997). Using the PA technique and electron paramagnetic resonance spectroscopy of nitroxide spin probes, this study reveals that the loss of desiccation tolerance results from a sequence of events involving unregulated metabolism followed by the loss of membrane permeability. Subsequently, pea seeds were dried at room temperature under a dry air flow (approximately 3% relative humidity [RH]) for 24 h (fast drying), and cucumber seeds were dried in a sealed container at 33% RH achieved by a saturated K2CO3 solution (Bruggink and van der Toorn, 1995). To induce desiccation tolerance in germinated cucumber radicles, 72-h-imbibed seeds were sorted into four classes according to the length of their protruding radicles (1, 2, 3, and 4 mm) and incubated at 10°C for 7 d in a PEG 8,000 solution having a water potential of −1.5 MPa (Bruggink and van der Toorn, 1995). Following incubation, seeds were rinsed in distilled water before analysis. After drying at 33% RH, seeds were re-imbibed on wet filter paper at 25°C. Seeds exhibiting growing radicle tips were scored as desiccation tolerant. Moisture contents were assessed gravimetrically by comparing the sample weights before and after drying for 40 h at 96°C, and are expressed on a dry weight basis.

**Determination of Membrane Integrity**

The effect of dehydration on membrane permeability was determined upon rehydration using an electron spin resonance (ESR) spin probe technique (Golovina et al., 1997, 1998; Leprince et al., 1999). The technique is based on the differential permeability of an amphiphilic spin probe and the broadening agent K3Fe(CN)6, which does not permeate through intact membranes. At various intervals of drying, three to four isolated pea and cucumber radicles were weighed, then imbibed in either 1 mM 2,2,6,6-tetramethyl-1-piperidinyloxy (Tempo) or 1 mM 4-oxo-2,2,6,6-tetramethyl-1-piperidinyloxy (Tempon) solutions for periods of time varying between 10 min for hydrated samples to 1 h for dried ones. Following imbibition, 120 mM K3Fe(CN)6 was added and incubation was prolonged for a further 15 min. Samples were retrieved from the solution and inserted into an ESR capillary tube. Spectra were recorded on an ESR X-band spectrometer (ESP 300, Bruker Analytik, Reinstetten, Germany) with the settings described in Golovina et al. (1997). After recording, samples were retrieved to determine their moisture content. The membrane permeability was assessed from the ratio between the amplitude of the peak given by the spin probe molecules present in the lipid fraction (L) and the amplitude of the peak given by the spin probe molecules dissolved in the aqueous cytoplasm (W). When membrane integrity is lost, the ESR signal from the spin probe present in the aqueous cytoplasm is reduced to invisibility by the polar broadening agent K3Fe(CN)6, whereas the signal of the probe present in the lipid remains intact. Therefore, the L/W ratio is high when the plasma membrane is disrupted. A low ratio indicates that the membrane is intact, since K3Fe(CN)6 cannot penetrate into the cell and broadens the signal of the spin probe in the cytoplasm.

**Noninvasive Analysis of Acetaldehyde, Ethanol, and CO2 Produced during Drying**

Acetaldehyde, ethanol, and CO2 were analyzed in the gas phase with the PA flow-through setup described in...
detail in Zuckermann et al. (1997) and Harren and Reuss (1997). As shown in Figure 1, the setup included successively: (a) three 5-mL glass cuvettes with an inlet and outlet in which radicles of seeds were dried; (b) a two-stage cold trap to remove water vapor that interferes with the measurements of other gases—the first stage was a Peltier device set at −5°C and the second a reservoir containing a stable level of liquid N₂ to achieve −50°C around the tubing; (c) a line-tunable CO laser that excites gas molecules specifically according to their IR fingerprint absorption; and (d) three parallel resonators, each coupled to a sensitive microphone (PA cell) in which the concentration of the gases is sequentially measured based on an acoustic phenomenon (for the theory and applications of the technique, see Harren and Reuss, 1997; Zuckermann et al., 1997). To discriminate and measure acetaldehyde, ethanol and CO₂ in the gas phase, seven infrared frequencies were selected (Table I). Measurement of each of the gases was taken every 9 min, during which time the frequencies were positioned by the laser setup, measured, and recorded on a computer. Thirty to 60 radicles were excised from imbibing seeds and incubated for 1 h on wet filter paper before the measurement. Radicles were laid on a glass fritter inside the glass cuvettes of known weight. Two cuvettes containing samples were analyzed simultaneously, whereas an empty cuvette was used as the control background for the gas detection and analysis. The cuvettes were connected to the cooling trap and flushed at 2 L h⁻¹ with synthetic air (CO₂, 200 μL L⁻¹) or a mixture of O₂:N₂ (1:1) that was passed through a cartridge containing anhydrous granules of CaCl₂ (20-mL bed volume) before entering the cuvette. All gas mixtures and flows were regulated by mass flow controllers.

Calculation of Volatile Release Rates as a Function of Moisture Content during Drying

The rates (R) of gas released by the dehydrating radicles over time were calculated according to the following equation:

\[ R(t) = \left( [G(t) - G_{\text{min}}] \times F \right) / DW \]  

(1)

where \( G(t) \) is the gas concentration (nanoliters per liter) measured by the photo-acoustic detectors at time \( t \), \( G_{\text{min}} \) is the lowest gas concentration detected during the entire measuring period, \( F \) is the air flow rate (liters per hour), and \( DW \) is the sample dry weight. Following calculation, patterns of gas emission comparable to those shown in A and E of Figure 2 were obtained. To correct for the downward drift of the baseline signal over time, the following fitting and calculation procedures were performed. First, the decline of the baseline with time in the absence of sample can be modeled as a double-exponential decay using the following equation (Fig. 2, step 1):

\[ R(t) = a \times e^{-bt} + c \times e^{-dt} \]  

(2)

where \( R(t) \) is the release rate at time \( t \), \( a \) and \( c \) are amplitudes of the decays, and \( b \) and \( d \) are the respective rate constants.
The equation was fitted to the data obtained before adding the sample in the cuvette and to those obtained with the sample once its moisture content was lower than approximately 0.07 g/g, when no metabolism could be detected. The values of the baseline in the presence of the sample were then interpolated using Equation 2 and subtracted from the data (Fig. 2, A and E). Furthermore, the measuring time was also normalized to the onset of drying. Thus, the absolute rates of gas released by the samples as a function of drying time were obtained as shown in Figure 2, B and F.

In desiccation-intolerant tissues, an upsurge of acetaldehyde was observed after a few hours of drying (Fig. 2F). To determine the rates of acetaldehyde evolving from dehydrating radicles as a function of water content, Examples were taken from dehydrating pea radicles isolated after 24 h of imbibition (desiccation-tolerant stage [A–D]) and 72 h of imbibition (desiccation-intolerant stage [E–H]). Step 1 (A and E): Plots of acetaldehyde release rates are shown as a function of measuring time in the presence of the sample (○). High values of acetaldehyde were found before adding the samples (panels 1). They were attributed to contamination from previous experiments, which was flushed away before adding the samples. The curve is the fit to the open symbols and corresponds to the baseline. Step 2 (B and F): After subtracting the fitted curve (shown in step 1) from the data, the acetaldehyde release rates as a function of time of drying were obtained. The arrows (F) mark the data points corresponding to the onset and end point of the upsurge of acetaldehyde release rates in desiccation-intolerant radicles. A new curve was fitted to the data points excluding those in between the arrows. The fit corresponds to an exponential decay (see “Materials and Methods”). Step 3: The exponential decay obtained in step 2 was subtracted from the data points including those in between the arrows. As a result, an acetaldehyde peak can be observed in desiccation-intolerant tissues (G) but not in desiccation-tolerant radicles (C). Step 4 (D and H): The times of drying were converted to moisture content using quadratic equation fitting the relation between drying time and moisture content (not shown) determined in parallel to the PA measurements.

Table I. Laserlines, infrared frequencies, and corresponding absorption coefficients of water, CO$_2$, acetaldehyde, and ethanol diluted in N$_2$.

<table>
<thead>
<tr>
<th>Laserlines</th>
<th>Infrared Frequency</th>
<th>Absorption Coefficient</th>
<th>Water</th>
<th>CO$_2$</th>
<th>Acetaldehyde</th>
<th>Ethanol</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>cm$^{-1}$</td>
<td>atm$^{-1}$ cm$^{-1}$</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>V13J7</td>
<td>1,933.42651</td>
<td>0.0079</td>
<td>0.00442</td>
<td>0.082</td>
<td>0.4</td>
<td></td>
</tr>
<tr>
<td>V13J11</td>
<td>1,765.45984</td>
<td>0.0219</td>
<td>–</td>
<td>21</td>
<td>0.08</td>
<td></td>
</tr>
<tr>
<td>V24J11</td>
<td>1,493.81274</td>
<td>0.0309</td>
<td>–</td>
<td>0.7</td>
<td>0.64</td>
<td></td>
</tr>
<tr>
<td>V28J18</td>
<td>1,406.90735</td>
<td>0.014</td>
<td>2.61 $10^{-5}$</td>
<td>4.7</td>
<td>7</td>
<td></td>
</tr>
<tr>
<td>V29J18</td>
<td>1,382.8562</td>
<td>0.0117</td>
<td>0.11</td>
<td>4.2</td>
<td>6</td>
<td></td>
</tr>
<tr>
<td>V13J9</td>
<td>1,772.88965</td>
<td>2.57</td>
<td>–</td>
<td>10.4</td>
<td>–</td>
<td></td>
</tr>
<tr>
<td>V19J11</td>
<td>1,616.04053</td>
<td>0.399</td>
<td>–</td>
<td>0.3</td>
<td>–</td>
<td></td>
</tr>
</tbody>
</table>
calculate the total amount of acetaldehyde released during the upsurge, the area beneath the peak was estimated. To extract the peak from the background signal, we used a fitting procedure to model the kinetics of acetaldehyde release during drying similar to that used for correcting the baseline. The model and fit are based on those developed by Voesenek et al. (1993) to describe the kinetics of ethylene release during desubmergence of Rumex plants. Using a PA setup similar to ours, these authors demonstrated that an exponential release of gases corresponds to the rapid diffusion of gases that are trapped in the tissues prior to the onset of the measurement rather than a progressive decrease in synthesis rates. When the upsurge in acetaldehyde is ignored (Fig. 2F, arrows), the release rates from the dehydrating sample over the time of drying, \( Y(t) \), was modeled as a negative exponential function using the equation:

\[
Y(t) = a \times e^{-bt}
\]

where \( t \) is the time of drying, \( a \) is the amplitude of the decay, and \( b \) is the rate constant. Occasionally, a third-order polynomial regression was used instead of Equation 3 because it gave a better fit (Fig. 2B). From Equation 3, the values of acetaldehyde release rates corresponding to the exponential decay were interpolated below the peak. The fitted values were then subtracted from the observed data to obtain the peak, the area beneath which was calculated (Fig. 2, C and G).

The final mathematical operation that was introduced relates the kinetics of the release of CO\(_2\), ethanol, and the upsurge of acetaldehyde as a function of moisture content during drying. To determine the water content in dehydrating radicles during the gas analysis, the cuvettes were disconnected and weighed at regular intervals during the measurement without disturbing the gas flow. When the rates of CO\(_2\) and acetaldehyde release reached a constant low value, the samples were retrieved to determine the dry weight and moisture content. A quadratic polynomial was fitted to the data (data not shown) to obtain the full range of moisture content values as a function of the time of drying.

### Fourier Transform Infrared (FTIR) Spectroscopy

FTIR spectra were recorded on an IR spectrometer (model 1725, Perkin Elmer, Beaconsfield, UK) equipped with a liquid N\(_2\)-cooled mercury/cadmium/telluride detector and a microscope (Perkin-Elmer) as described previously (Wolkers and Hoekstra, 1995). Palmitoyloleoylphosphatidylcholine (POPC) vesicles were prepared in the presence or absence of acetaldehyde, as outlined in Wolkers et al. (1999). Samples (5 \( \mu \)L) were dried directly on circular CaF\(_2\) windows (13 \( \times \) 2 mm) for at least 3 h in a stream of dry air at 23°C (<3% RH). Before the samples were removed from the dry-air box, another window was placed on top of the sample window with a rubber ring in between, and then mounted in a temperature-controlled brass cell. Hydrated vesicle samples were concentrated by ultracentrifugation, and the pellet was used for FTIR analysis. After cooling to \(-50^\circ\text{C}\), the sample was heated at a scanning rate of 1°C/min to 80°C, while spectra were recorded every 1.5 min. Phase transitions were derived from wave number-temperature plots following procedures described in Wolkers et al. (1999).

### Experimental Design and Statistical Treatment

Tolerance of desiccation was tested at least three times on 30 to 50 seeds per treatment. For membrane permeability analysis, the experimental unit was a sample of three to four isolated radicles dried to a certain moisture content. Experiments were repeated until a range of moisture contents was covered, from fresh to dried material. The gas analysis during drying was arranged in a randomized complete block design (four to six replicates, three stages of desiccation tolerance, and three measuring cuvettes). A single drying experiment of 30 to 60 radicles was considered as the experimental unit for each species. A Student’s \( t \) test was used to compare the effect of drying and atmosphere composition on the upsurge of acetaldehyde. Data were fitted with polynomial or exponential functions using the Marquardt-Levenberg algorithm (SigmaPlot, Jandel Software, San Rafael, CA). The fitting accuracy was systematically gauged by the parameter coefficients of variation that ranged between 4% and 10%. The adjusted \( r^2 \) values of the fitted curves were always >0.95.

### RESULTS

#### Desiccation Tolerance and Membrane Integrity in Germinating Pea and Cucumber Radicles

After 42 h of imbibition at 20°C, cucumber seeds had not started to germinate and were still desiccation tolerant. This was confirmed using an ESR spin probe technique to assess the integrity of plasma membranes upon rehydration after different periods of drying (Golovina et al., 1997). Figure 3B shows that membrane permeability in 42-h-old imbibed radicles remained low throughout drying. After 72 h of imbibition, 97% of the seeds had germinated. They exhibited a radicle length ranging between 0.5 and 4 mm. Radicles exhibiting a length longer than 2 mm were not able to survive drying (Fig. 3A). However, they could be dried to 0.29 g/g without affecting the membrane permeability (Fig. 3B; Table II). Upon further drying, the permeability increased with decreasing moisture content. Thus, a critical moisture content below which membranes become sensitive to drying can be determined from the breakpoint in the relations between decreasing moisture content and membrane permeability (Table II).

Exposing germinated seeds to a mild water stress reinduced desiccation tolerance in sensitive radicles of several species (Bruggink and van der Toorn, 1995). In this study, desiccation tolerance was reinduced in 72-h-old imbibed radicles of cucumber by incubating seeds in a PEG solution equivalent to a water potential of \(-1.5\) MPa at 10°C for 7 d prior to dehydration (Fig. 3A). Furthermore, the plasma membranes of PEG-treated radicles remained...
intact throughout drying and did not exhibit a critical moisture content (Fig. 3B).

In pea, 24- and 72-h-old imbibed radicles were chosen to represent the desiccation-tolerant and -intolerant stages, respectively. During dehydration of 24-h-old radicles the plasma membrane permeability remained unchanged (Fig. 3C). In contrast, 72-h-old radicles started to lose their plasma membrane integrity when the tissues were dried below 1.2 g/g (Fig. 3C; Table II).

Response of CO₂ Production to Drying

Rates of CO₂ production from drying radicles were analyzed to assess whether there is a repression of respiration associated with desiccation tolerance. Preliminary experiments showed that incubating isolated radicles on wet filter paper did not induce significant changes in the CO₂ emission rates over a period of 4 h (data not shown). Drying induced a progressive decrease in rates of CO₂ emission as the radicles of both cucumber and pea lost water (Fig. 4). In both species, the rates of CO₂ produced by desiccation-sensitive radicles during dehydration were at least 2-fold higher than those in tolerant radicles. In cucumber the incubation in PEG at -1.5 MPa that induces desiccation tolerance of germinated radicles was found to drastically reduce the respiration rates (Fig. 4B). Before drying, the CO₂ emission rates in PEG-treated radicles were 5.1-fold lower than in untreated ones. During dehydration, rates of CO₂ emission in PEG-treated radicles remained low and steady until around 1 g/g, and thereafter decreased rapidly (Fig. 4B). In desiccation-sensitive tissues of both species, the decline in CO₂ emission rates was less pronounced at the onset of drying than below approximately 2.5 g/g. However, the progressive loss of membrane integrity during drying did not clearly affect CO₂ emission. In desiccation-intolerant pea radicles, the responses of CO₂ emission rates to drying were similar whether the tissues were dried in air or in 50% O₂ (data not shown). It was not possible to estimate the moisture content at which the CO₂ emission ceased during drying because of the large errors made with the quadratic polynomial equation fitting the relation between drying time and dehydration.

Kinetics of Emission of Acetaldehyde and Ethanol during Drying

It has been found that the rise in cytoplasmic viscosity and the resulting impeded diffusion of O₂ during drying imposed hypoxia-like conditions on the metabolism of desiccation-tolerant seeds and pollen (Leprince and Hoekstra, 1998). This prompted us to examine whether the kinetics of release of volatiles from alcoholic fermentation during dehydration of pea and cucumber radicles are linked with desiccation tolerance. After correction for the background, the typical pattern of acetaldehyde production from dehydrating radicles is shown in Figure 2, B and F. At time 0 of drying, isolated radicles were found to produce large amounts of acetaldehyde, regardless of the level of desiccation tolerance. Furthermore, after 1 h of incubation on wet filter paper, the acetaldehyde production rates of isolated radicles varied between 0.5 and 5 μL h⁻¹ g⁻¹ dry weight in cucumber and between 2 and 40 μL h⁻¹ g⁻¹ dry weight in pea. When data were expressed as a function of drying time, the kinetics of acetaldehyde release during dehydration fitted an exponential decay (Fig. 2F). In PA measurements, such decay is due to the progres-
sive release of gases that were trapped in the tissues before drying rather than a decline of synthesis rates (see "Materials and Methods," Voesenek et al., 1993). It is likely that during dehydration, the removal of the apoplastic water facilitates the diffusion of gases throughout the tissues. This interpretation is further supported by PA measurements showing that water vapor released during drying also followed an exponential decay (data not shown).

During drying, the exponential decay was interrupted by an upsurge in the acetaldehyde release rates, indicating that drying induced a synthesis of acetaldehyde. This upsurge was found solely in dehydrating desiccation-intolerant tissues of pea and cucumber. To characterize this upsurge, the peak was subtracted from the exponential decay (Fig. 2G) and plotted as a function of moisture content during drying (Fig. 2H). The procedure was performed with the acetaldehyde release measurements in all treatments (Fig. 5). The area under the peak and the moisture content corresponding to the amplitude of the peak were reported in Table II. Figure 5 shows that in dehydrating desiccation-sensitive tissues, the acetaldehyde production started at 3.6 and 3.3 g/g in pea and cucumber radicles, respectively. In the presence of 50% O2, the acetaldehyde upsurge from 72-h-imbibed pea radicles was 3-fold lower than that of tissues dried in air, indicating that acetaldehyde emission during drying is linked to hypoxic conditions (Table II). In cucumber radicles, the desiccation-induced acetaldehyde peak was 7.3-fold lower in the PEG-treated radicles than in the untreated 72-h-imbibed tissues (Table II). The production rates were maximum at 1.73 and 0.94 g/g in desiccation-intolerant pea and cucumber radicles, respectively, a value much higher than the critical moisture content at which the membranes of the respective species started to be disrupted by the drying treatment (Table II). Therefore, we conclude that drying induces disturbances in metabolism associated with the desiccation sensitivity of the tissue well before the loss of membrane integrity.

The kinetics of ethanol release during drying could not be thoroughly analyzed because of an approximately 30-min delay in the response of the PA detection of ethanol, as illustrated in dehydrating desiccation-tolerant pea radicles (Fig. 6A, dashed lines). Nevertheless, the plots of ethanol release rates versus time of drying show different kinetics in dehydrating desiccation-tolerant and -intolerant radicles of pea. In desiccation-tolerant tissues, the kinetics of ethanol emission followed an exponential decay similar to that found for acetaldehyde (Fig. 6A), suggesting that no net synthesis of ethanol occurred during drying. In desiccation-
sensitive radicles, the ethanol release rates were low and remained unchanged during the first hour of drying. After 2 h of drying and onward, desiccation-sensitive radicles were found to accelerate the production of ethanol. When data were plotted as a function of moisture content during drying, the upsurge of ethanol occurred when the tissues were dried below 2 g/g, which approximately coincided with the acetaldehyde peak (Fig. 6B). In dehydrating cucumber radicles, the ethanol release rates followed an exponential decay without upsurge, regardless of their level of desiccation tolerance (data not shown).

Effect of Acetaldehyde on POPC Phase Behavior and Desiccation Tolerance

Loss of membrane permeability is linked with changes in the fluidity and phase properties of membranes (Senaratna et al., 1984, 1987; Van Bilsen et al., 1994; Hoekstra et al., 1997). To gain insight into the effects of acetaldehyde on the properties of phospholipid bilayers, the band positions of the symmetric CH$_2$ stretching vibration of POPC vesicles in the presence and absence of acetaldehyde was obtained from the FTIR spectra and plotted against temperature (Fig. 7). Low amounts of acetaldehyde (mass ratio 1:1.3 acetaldehyde:POPC before drying) were found to have two effects on the POPC bilayer properties. First, in the hydrated state, acetaldehyde slightly depressed the gel- to liquid-crystalline phase transition temperature ($T_m$) of POPC vesicles. In the dried state, the depression of $T_m$ by acetaldehyde was more pronounced (Fig. 7). Second, below the $T_m$, the presence of acetaldehyde in the hydrated POPC vesicles induced a slight upward shift in wave number, indicating that the CH$_2$ groups have more vibrational freedom. In the dried state, the same effect was observed, indicating that acetaldehyde exerts disordering effects on the acyl chains in the gel phase at room temperature.

Figure 5. The effect of drying on the upsurge in the acetaldehyde release rates in imbibed radicles of pea (A) and cucumber (B). In pea, desiccation-intolerant tissues were dried under air (○) or 50% O$_2$ (●), whereas desiccation-tolerant radicles (○) were dried under air. In cucumber, all experiments were performed in air using desiccation-intolerant (●), desiccation-tolerant (○) and PEG-induced desiccation-tolerant (▲) radicles. For each treatment, one representative experiment is shown. The arrows indicate the critical moisture content corresponding to the onset of membrane damage.

Figure 6. The effect of drying on the ethanol release rates from imbibed radicles of pea at the desiccation-tolerant (○) and desiccation-intolerant stages (●). Data were plotted as a function of time of drying (A) and water content (B). The dashed lines and the arrow in panel A mark the lag in the time between the start of the drying experiment (and thereby ethanol emission by the tissues) and the ethanol detection by the PA cells. The lag time was always present during our measurements. It was clearly visible in dehydrating desiccation-tolerant tissues. The reason for the lag time was not investigated. In B, the data on desiccation-intolerant radicles (●) are shown only to demonstrate the occurrence of the ethanol upsurge at a moisture content similar to that of acetaldehyde (Fig. 5A). The moisture contents were obtained as described in Figure 2.
show that a balanced down-regulation leading to metabolic arrest during dehydration is associated with desiccation tolerance of seeds. Using germinating pea and cucumber radicles, we compared the kinetics of CO\textsubscript{2} emission as a marker for respiration and energy metabolism and the kinetics of fermentation as a marker of imbalanced metabolism in response to dehydration of desiccation-tolerant and -intolerant tissues. For this purpose, it was necessary to use a PA technique to detect minute changes in gas release rates without disturbing the dehydration process. Furthermore, we took advantage of the amenability of PEG treatments, which can reinduce desiccation tolerance in germinated cucumber radicles (Bruggink and van der Toorn, 1995).

### Reduction of CO\textsubscript{2} Production Associated with Desiccation Tolerance

In both pea and cucumber, there was no clear evidence that dehydration provokes a depression of CO\textsubscript{2} production in desiccation-tolerant radicles. Nevertheless, CO\textsubscript{2} production during drying was 2-fold higher in desiccation-intolerant compared with desiccation-tolerant radicles. Thus, at a similar moisture content during drying, the production and demand for energy appears to be higher in desiccation-intolerant tissues than in desiccation-tolerant tissues. Interestingly, incubation of 72-h-old imbibed radicles in PEG induced an important decrease in CO\textsubscript{2} emission rates compared with untreated 72-h-old radicles at time 0 of drying (Fig. 4). This decrease in CO\textsubscript{2} production occurred in conjunction with the induction of desiccation tolerance, suggesting that energy metabolism must be lowered before the tissues start to lose water in order to achieve desiccation tolerance. Apparently, the repression of CO\textsubscript{2} production is not induced during dehydration but must be present before the tissues lose water.

Since desiccation tolerance is a multifactorial trait (Leprine et al., 1993; Vertucci and Farrant, 1995), there is always the possibility that reduction of metabolites in...
Dehydration of Desiccation-Sensitive Germinating Radicles Results in Imbalanced Metabolism

An important requirement for metabolic arrest is that rates of all cellular processes be reduced in concert so that metabolism is rebalanced at much lower ATP costs (Hand and Hardewig, 1996). If the down-regulation leading to total metabolic arrest is not well coordinated, imbalance in metabolic rates may result in the accumulation of by-products to toxic levels. If imbalanced metabolism is associated with desiccation sensitivity in seeds, it should be observed before the loss of membrane integrity. Otherwise, it can be argued that the loss of metabolic control is the consequence of membrane damage (Kimmerer and Kozlowski, 1982). Our non-invasive analysis of fermentation products (Figs. 5 and 6), in conjunction with the assessment of membrane damage during dehydration (Fig. 3), provides evidence that the desiccation tolerance is associated with a tight control of metabolism during dehydration. Desiccation-sensitive radicles of pea were found to produce acetaldehyde and ethanol during drying, whereas in cucumber radicles, an upsurge was found for acetaldehyde only (Fig. 5; Table II). These upsurges occurred at different moisture contents in pea and cucumber and preceded the only (Fig. 5; Table II). These upsurges occurred at different moisture contents in pea and cucumber and preceded the loss of membrane integrity. Otherwise, it can be argued that the loss of metabolic control is the consequence of membrane damage (Kimmerer and Kozlowski, 1982). Our non-invasive analysis of fermentation products (Figs. 5 and 6), in conjunction with the assessment of membrane damage during dehydration (Fig. 3), provides evidence that the desiccation tolerance is associated with a tight control of metabolism during dehydration. Desiccation-sensitive radicles of pea were found to produce acetaldehyde and ethanol during drying, whereas in cucumber radicles, an upsurge was found for acetaldehyde only (Fig. 5; Table II). These upsurges occurred at different moisture contents in pea and cucumber and preceded the loss of membrane integrity. Otherwise, it can be argued that the loss of metabolic control is the consequence of membrane damage (Kimmerer and Kozlowski, 1982). Our non-invasive analysis of fermentation products (Figs. 5 and 6), in conjunction with the assessment of membrane damage during dehydration (Fig. 3), provides evidence that the desiccation tolerance is associated with a tight control of metabolism during dehydration. Desiccation-sensitive radicles of pea were found to produce acetaldehyde and ethanol during drying, whereas in cucumber radicles, an upsurge was found for acetaldehyde only (Fig. 5; Table II). These upsurges occurred at different moisture contents in pea and cucumber and preceded the loss of membrane integrity. Otherwise, it can be argued that the loss of metabolic control is the consequence of membrane damage (Kimmerer and Kozlowski, 1982).
Is Acetaldehyde Production Responsible for Membrane Damage during Drying?

This study shows that in desiccation-sensitive tissues, drying results in an increased production of acetaldehyde before the onset of membrane damage. Wounding, ozone, temperature, and water stress in seedlings of several species have been found to induce the synthesis of large amounts of acetaldehyde and ethanol (Kimmerer and Kozlowski, 1982; Conley et al., 1999). The cytotoxicity of acetaldehyde also has been invoked to explain the cytoplasmic male sterility in pollen and degeneration of the tapetum layer (Tadege and Kuhlemeier, 1997) and loss of seed viability during storage (Zhang et al., 1997). Figure 7 clearly shows that acetaldehyde at physiological concentrations may partition into the phospholipids and alter the phase properties of membranes. Increased membrane permeability upon rehydration of dried systems has been correlated both with the partitioning of amphiphilic substances (Golovina et al., 1998) and alcohols (Priestley and Leopold, 1980) into phospholipids and with changes in membrane phase behavior (Senaratna et al., 1984; Van Bilsen et al., 1994). In mammals, acetaldehyde is known to favor lipid peroxidation (Lieber, 1988), a factor that has been correlated with the loss of desiccation tolerance in seeds (Senaratna et al., 1987; Hendry et al., 1992; Leprince et al., 1993, 1994). In the light of these data and those shown in Figure 5, it is tempting to link the synthesis of acetaldehyde during drying with the loss of membrane integrity. Incubating cucumber seeds in a non-lethal concentration of acetaldehyde had no significant effects on desiccation tolerance (Table III). Our data are in contradiction with both the literature on acetaldehyde toxicity in mammals (Lieber, 1988) and the observation made by Zhang et al. (1997) on lettuce seeds. In cucumber seeds, we cannot exclude the possibility that acetaldehyde did not reach the sites of action in the membranes because it did not permeate into the tissues, because it escaped during drying, or because it was metabolized. Thus, bearing in mind the toxicity of acetaldehyde on seeds and mammals (Lieber, 1988; Zhang et al., 1997), its reactivity toward lipids, proteins, and nucleosides (Fraenkel-Conrat and Singer, 1988; Lieber, 1988), and its perturbing effect on membrane permeability and significant for desiccation tolerance. Plant Physiol 118: 975–986


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