Influence of Water Content and Temperature on Molecular Mobility and Intracellular Glasses in Seeds and Pollen

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Although the occurrence of intracellular glasses in seeds and pollen has been established, physical properties such as rotational correlation times and viscosity have not been studied extensively. Using electron paramagnetic resonance spectroscopy, we examined changes in the molecular mobility of the hydrophilic nitroxide spin probe 3-carboxy-proxyl during melting of intracellular glasses in axes of pea (Pisum sativum L.) seeds and cattail (Typha latifolia L.) pollen. The rotational correlation time of the spin probe in intracellular glasses of both organisms was approximately $10^{-3}$ s. Using the distance between the outer extrema of the electron paramagnetic resonance spectrum ($2\Delta\tau_x$) as a measure of molecular mobility, we found a sharp increase in mobility at a definite temperature during heating. This temperature increased with decreasing water content of the samples. Differential scanning calorimetry data on these samples indicated that this sharp increase corresponded to melting of the glassy matrix. Molecular mobility was found to be inversely correlated with storage stability. With decreasing water content, the molecular mobility reached a minimum, and increased again at very low water content. Minimum mobility and maximum storage stability occurred at a similar water content. This correlation suggests that storage stability might be at least partially controlled by molecular mobility. At low temperatures, when storage longevity cannot be determined on a realistic time scale, $2\Delta\tau_x$ measurements can provide an estimate of the optimum storage conditions.

Deterioration of seeds and pollen during storage involves many physical and chemical changes, such as disrupted intracellular integrity, decreased activities of enzymes, lipid peroxidation and deesterification, and Maillard reactions (Priestley, 1986; Wilson and McDonald, 1986; Wettlaufer and Leopold, 1991; Van Bilsen and Hoekstra, 1993; Van Bilsen et al., 1994). Since the formation of glasses in dehydrating biological tissues has been established, this physical phenomenon has been put forward as a prominent factor in the control of deterioration rates during storage (Burke, 1986; Williams and Leopold, 1989; Leopold et al., 1994; Leprince and Walters-Vertucci, 1995; Buitink et al., 1996). A glass is a thermodynamically unstable solid-state with an extremely high viscosity (Franks et al., 1991). Its formation is promoted by low water content of tissues and by low temperatures. Both factors are also known to extend the longevity of seeds and pollen (Roberts, 1972; Roberts and Ellis, 1989; Vertucci and Roos, 1993; Buitink et al., 1998), and improved storage stability was observed when glasses were present (Sun and Leopold, 1994; Sun, 1997; Buitink et al., 1998). It is assumed that the high viscosity of intracellular glasses decreases molecular mobility and impedes diffusion within the cytoplasm, thus slowing down deleterious reactions and changes in structure and chemical composition during aging (Sun and Leopold, 1993; Sun, 1997). However, the molecular mobility and viscosity in biological glasses has received little attention.

Molecular mobility has been studied using EPR spectroscopy by labeling polymers and food materials with a suitable, stable spin probe (Steffen et al., 1992; Blackburn et al., 1996; Dzuba, 1996; Hemminga and Van den Dries, 1998). From the EPR spectra of the spin probe, $\tau_R$ can be assessed (Kumler and Boyer, 1976; Kovarskii et al., 1978; Spielberg and Gelerinter, 1982; Ohta and Kuwata, 1985; Roozen and Hemminga, 1990; Roozen et al., 1991; Dzuba et al., 1993). Whereas $\tau_R$ values of $10^{-12}$ to $10^{-9}$ s can be calculated from conventional EPR spectra (Knowles et al., 1976), ST-EPR spectroscopy further expands this range to very slow ($10^{-9}$–$10^{-3}$ s; Hemminga, 1983) and ultra slow ($10^{-3}$–$10^{-2}$ s; Van den Dries et al., 1998) molecular motions. ST-EPR spectroscopy has been successfully applied to determine $\tau_R$ values of spin probes in sugar glasses (Roozen et al., 1991; Van den Dries et al., 1998) and organic liquids at low temperatures (Ito, 1983). In glassy Suc-water and malt-o-oligosaccharide mixtures, $\tau_R$ decreases by several orders of magnitude upon approaching $T_g$ (Roozen et al., 1991).

The present paper is aimed at gaining insight into changes in the molecular mobility that accompany glass formation in anhydrobiotes. We used EPR and ST-EPR spectroscopy to characterize the molecular motion of CP, the polar nitroxide spin probe that we incorporated into axes of pea (Pisum sativum L.) seeds and cattail (Typha latifolia L.) pollen. We show that the distance between the

Abbreviations: CP, 3-carboxy-proxyl; EPR, electron paramagnetic resonance; ST-EPR, saturation-transfer EPR; $\tau_R$, rotational correlation time; TEMPO, 2,2,6,6-tetramethyl-1-piperidinylkoxy; $T_g$, glass-to-liquid transition temperature.
EPR spectroscopy can be used to detect changes in molecular motion, and that ST-EPR spectroscopy allows an estimation of $\tau_R$. We discuss the possible relationship between glasses, molecular mobility, and storage stability in these organisms.

MATERIALS AND METHODS

Plant Material and Treatments

Mature male inflorescences of cattail (Typha latifolia L.) were collected from field populations near Wageningen, The Netherlands, in 1996, and allowed to shed their pollen in the laboratory. Pollen (94% germination) was cleaned by sieving through a fine copper mesh, dried in dry air to 0.05 g water g$^{-1}$ dry weight, and stored at -20°C until use. Pea (Pisum sativum cv Carina) seeds (99% germination) were obtained from Nunhems Zaden (Haelen, The Netherlands) and stored at 5°C until use.

The polar nitroxide spin probe CP (Sigma) was used for spin labeling of pea seeds and cattail pollen. Pollen (3 g) was prehydrated in water vapor for 16 h at 5°C to about 0.7 g water g$^{-1}$ dry weight, and then mixed at 25°C with 6 mL of liquid germination medium containing 2.5 mm CP. The germination medium consisted of 1.6 mm H$_3$BO$_3$, 1.3 mm Ca(NO$_3$)$_2$·4H$_2$O, 0.8 mm MgSO$_4$·7H$_2$O, 1.0 mm KNO$_3$, and 0.2 mm Suc in 2 mm sodium-phosphate-citrate buffer, pH 5.9. After a few minutes, an additional 20 mL of the germination medium was added, and the pollen was recovered by filtration. The pollen was then mixed with 20 mL of a solution of 1 mm CP and 120 mm of the broadening agent potassium ferricyanide. Ferricyanide broadens spin-probe signals in the solution surrounding the cells to invisibility (Golovina and Tikhonov, 1994; Golovina et al., 1997). After 5 more min, the pollen was recovered by filtration, spread out in a large Petri dish, and rapidly dried in a flow of dry nitrogen vapor as the coolant. Samples were rapidly cooled to -150°C and allowed to equilibrate for 30 min, after which scans were recorded at 10°C increments with equilibration for 5 min after each increment.

Conventional EPR spectroscopy can detect changes in $\tau_R$ of spin probes ranging from $10^{-12}$ to $10^{-9}$ s, which corresponds to the lifetime of the probe in a given orientation. In this motional range, the EPR spectrum of nitroxides consists of three lines (Fig. 1, top spectrum), and $\tau_R$ can be calculated according to the method of Knowles et al. (1976):

$$\tau_R = 6.5 \times 10^{-10} \Delta B_0 [(h_0 / h_{-1})^{1/2} - 1]$$

(Eq. 1)

![Figure 1. EPR spectra of CP in cattail pollen at various water contents (g/g, g water g$^{-1}$ dry weight). Spectra were recorded at room temperature. The $h_{-1}$, $h_0$, and $h_1$ peaks are shown in the top spectrum. The distance between the two outer extrema, $2A_{\Delta \tau}$, is indicated in the bottom spectrum.)

EPR and ST-EPR Measurements

EPR spectra were recorded with an X-band EPR spectrometer (model 300E, Bruker Analytik, Rheinstetten, Germany). Microwave power was kept low (200 $\mu$W or 2 mW) to avoid saturation. Modulation amplitude was 0.4 G for pea axes and 1 G for cattail pollen.

Samples with various water contents were loaded into a 3-mm-diameter EPR capillary. For each measurement, the capillary was filled for a length of 5 cm with pollen or with two isolated pea axes. To prevent water loss during the measurements, the capillaries were sealed at both sides. Temperature was controlled using a controller with liquid nitrogen vapor as the coolant. Samples were rapidly cooled to -150°C and allowed to equilibrate for 30 min, after which scans were recorded at 10°C increments with equilibration for 5 min after each increment.

Figure 1. EPR spectra of CP in cattail pollen at various water contents (g/g, g water g$^{-1}$ dry weight). Spectra were recorded at room temperature. The $h_{-1}$, $h_0$, and $h_1$ peaks are shown in the top spectrum. The distance between the two outer extrema, $2A_{\Delta \tau}$, is indicated in the bottom spectrum.
where \( h_{-1} \) and \( h_0 \) are the heights of the high-field and central lines in the EPR spectra, respectively, and \( \Delta B_0 \) is the line width of the central line in Gauss. The rotational motion of the spin probe was assumed to be isotropic. The \( \tau_R \) of partially hydrated (above approximately 0.25 g water g\(^{-1}\) dry weight) pollen and pea axes was determined using Equation 1.

The characteristic \( \tau_R \) for spin probes in organic glasses near and above \( T_g \) is approximately \( 10^{-5} \) s or higher (Dzuba et al., 1984; Roozen et al., 1991; Van den Dries et al., 1998). These \( \tau_R \) values cannot be determined using Equation 1, because the shapes of the lines in the spectra change due to the appearance of a powder spectrum (Fig. 1, below 0.25 g water g\(^{-1}\) dry weight). In these powder spectra the distance between the two outer extrema (2A\(_{zz}\)) is temperature dependent for a number of glass-forming substances at and above their \( T_g \) (Kumler and Boyer, 1976; Kovarskii et al., 1978; Spielberg and Gelernter, 1982; Ohta and Kuwata, 1985; Dzuba et al., 1993) (Fig. 1). We made use of this parameter to obtain a qualitative measurement of the molecular mobility of CP.

From the \( \tau_R \) one can derive the viscosity of the matrix in which the spin probe is rotating, according to the modified Stokes-Einstein equation (Roozen et al., 1991):

\[
\tau_R = \left( \eta V/k_b T \right) k + \tau_0
\]

where \( \eta \) is the solvent viscosity, \( k_b \) is Boltzmann's constant, \( V \) is the volume of the rotating molecule, \( T \) is the absolute temperature, \( \tau_0 \) is the zero viscosity \( \tau_R \), and \( k \) is a dimensionless slip parameter.

ST-EPR spectroscopy was used in the motional region for \( \tau_R > 10^{-7} \) s. This method is based on the diffusion and recovery of saturation between different portions of the powder spectrum in competition with field modulatio (Hemminga, 1983). For ST-EPR spectroscopic measurements, the second harmonic quadrature absorption signal was detected under the following conditions: a field-modulation amplitude of 5 G, microwave power of 100 mW, and field-modulation frequency of 50 kHz (Hemminga et al., 1984). The phase was set with the self-null method (Thomas et al., 1976).

In ST-EPR spectroscopy, \( \tau_R \) values are usually obtained empirically using reference material with known viscosity. Here, spectra of CP in anhydrous glycerol were used to construct a calibration curve according to the method of Van den Dries et al. (1998). Because the viscosity for anhydrous glycerol is known over a broad temperature range, \( \tau_R \) of CP in glycerol can be obtained from Equation 2. Spectra of CP in anhydrous glycerol were recorded every 3°C, and the values of the line-shape parameters \( L^*/L \) and \( C^*/C \) (explained in Fig. 5) were calculated for each temperature (data not shown). From the curves representing the line-shape parameters of CP in glycerol against \( \tau_R \), the \( \tau_R \) values of CP in the axes and pollen were obtained by interpolation of the corresponding line-shape parameters.

### Differential Scanning Calorimetry

Pollen and pea axes with different water contents were hermetically sealed in aluminum pans for differential scanning calorimetry. Second-order transitions of the samples were determined using a differential scanning calorimeter (Pyris 1, Perkin-Elmer) calibrated for temperature with indium (156.6°C) and methylene chloride (−95°C) standards and for energy with indium (28.54 J g\(^{-1}\)). Baselines were determined using an empty pan, and all thermograms were baseline corrected. Scans were taken from −100°C to 120°C at a rate of 10°C min\(^{-1}\). The \( T_m \) values were determined by the onset and midpoint of the temperature range over which the change in specific heat occurred. All analyses were performed with Perkin-Elmer software.

### RESULTS

#### Molecular Motion in Pollen and Seeds

Figure 1 shows representative EPR spectra of CP in cattail pollen at different water contents recorded at room temperature. When the water content was decreased from 0.53 to 0.25 g water g\(^{-1}\) dry weight (top two spectra), the relative amplitudes of the outer spectral lines decreased and the width of the central peak increased. Using Equation 1, the \( \tau_R \) of CP in hydrated pollen and pea axes (approximately 1.2 g water g\(^{-1}\) dry weight) was calculated to be approximately \( 10^{-11} \) s (data not shown). In the partially dehydrated state (0.53 g water g\(^{-1}\) dry weight), \( \tau_R \) of CP in cattail pollen was calculated as \( 1.6 \times 10^{-10} \) s (Fig. 1, top spectrum). At 0.25 g water g\(^{-1}\) dry weight (Fig. 1, second spectrum from the top), \( \tau_R \) of CP in pollen was 7.9 \( \times 10^{-10} \) s. The \( \tau_R \) of CP in pea axes at 0.5 g water g\(^{-1}\) dry weight was 3.6 \( \times 10^{-10} \) s (data not shown). At a water content below 0.25 g water g\(^{-1}\) dry weight, a powder spectrum (characterized by the two broad peaks at the extremes) overlapped the mobile spectrum (three sharp lines separated by a distance of 15 G). The bottom spectrum shown in Figure 1 has the characteristic shape of a powder spectrum, indicative of slow molecular mobility of the spin probe with \( \tau_R > 10^{-8} \) s. With the appearance of this powder spectrum below 0.25 g water g\(^{-1}\) dry weight, the \( \tau_R \) of CP cannot be calculated using Equation 1, because the line shapes are distorted. The distance between the two broad peaks at the extremes is referred to as 2A\(_{zz}\) (Fig. 1, bottom spectrum). Similar spectra were obtained for pea axes in relation to water content (data not shown).

Figure 2 shows EPR spectra of CP in dry pea axes and cattail pollen (both having 0.07 g water g\(^{-1}\) dry weight) at a range of temperatures. At least two overlapping spectra contributed to the total spectrum observed: a powder spectrum at all temperatures and a mobile spectrum above 20°C, the contribution of which increased with increasing temperature. Note that the contribution of the mobile component to the total spectrum is considerably larger for pollen at 70°C than for pea axes at 90°C.

At −150°C, we assumed that the motion of the probe was completely immobilized and therefore we took the corresponding 2A\(_{zz}\) values as the maximum values. At this low temperature, the value of 2A\(_{zz}\) gives information about the polarity of the spin probe’s environment in the tissue (Knowles et al., 1976). In pea axes the maximum 2A\(_{zz}\)
decreased with decreasing water content from 74 to 70 G, whereas in cattail pollen, it changed from 72.5 to 71.5 G (Fig. 3).

Because we observed a powder spectrum of CP in pea axes and cattail pollen at low water contents (<0.2 g water g\(^{-1}\) dry weight), \(\tau_R\) cannot be directly calculated from the EPR spectra using Equation 1. However, the change in \(2A_{zz}\) with temperature can be used as an estimate of molecular motion (Van et al., 1974; Dzuba, 1996). A decrease in \(2A_{zz}\) is indicative of an increase in molecular mobility. Figure 4 shows these changes in \(2A_{zz}\) with temperature. When the temperature of pea axes and cattail pollen increased, the \(2A_{zz}\) slowly decreased, then abruptly decreased above a definite temperature. With increasing water contents, this abrupt decrease in \(2A_{zz}\), denoting an abrupt increase in molecular mobility, commenced at lower temperatures. At a water content of 0.002 g water g\(^{-1}\) dry weight in pea axes the decrease in \(2A_{zz}\) was less clear.

We used ST-EPR to obtain an indication of the magnitude of change in \(\tau_R\) (reflecting the lifetime of the probe in a given orientation) that corresponds to the change in \(2A_{zz}\) with temperature. This technique is based on recording spectra under saturation conditions, which yields line shapes that are sensitive to \(\tau_R > 10^{-7}\) s (Hyde and Dalton, 1979). The ST-EPR spectra can be well characterized by independent line-shape parameters such as the line-height ratios L'/L' and C'/C (Fig. 5). These ratios are dependent on the \(\tau_R\) of the spin probe. To determine the \(\tau_R\) of CP corresponding to a certain line-height ratio, anhydrous

**Figure 2.** EPR spectra of CP in pea axes and cattail pollen recorded at various temperatures. Both contained approximately 0.07 g water g\(^{-1}\) dry weight.

**Figure 3.** Change in the distance between the outer extrema of the EPR spectra (\(2A_{zz}\)) of CP in pea axes (○) and cattail pollen (●) at −150°C as a function of water content. dw, Dry weight.

**Figure 4.** Comparison of the distance between the outer extrema of the EPR spectra (\(2A_{zz}\)) of CP in pea axes and cattail pollen at different water contents against temperature.
glycerol was used as a reference solvent (this solvent was previously used as a reference in sugar glasses in Roozen et al. [1991] and Van den Dries et al. [1998]). Although the composition of the cytoplasm in seeds and pollen is not comparable to glycerol, at present the use of glycerol will give the best approximation of the relationship between the line-height ratios and \( t_R \). Extrapolation of the \( t_R \) of CP in seeds and pollen to the corresponding viscosity is not valid, because the Stokes-Einstein law fails at temperatures below approximately 1.3 \( T_g \) (Liu and Oppenheim, 1996).

As the viscosity for glycerol is known over a broad temperature range, the corresponding \( t_R \) could be calculated according to Equation 2. Subsequently, the two line-height ratios were obtained from ST-EPR scans of anhydrous glycerol for a range of temperatures and plotted against the corresponding \( t_R \). Those calibration curves were used to obtain \( t_R \) from line-height ratios calculated from ST-EPR scans of CP in pea axes (Fig. 5). The \( t_R \) values for pea axes (0.08 g water g\(^{-1}\) dry weight) calculated according to both line-height ratios and \( t_R \) derived from the \( C/C \) ratio show an increase in mobility with increasing temperature comparable to the 2A\(_{zz}\) curves (Fig. 4).

One can speculate on what causes the difference in \( t_R \) obtained from both line-height ratios. The \( t_R \) obtained from the \( L''/L' \) ratio probably reflects overall isotropic rotational motion, whereas the \( t_R \) derived from \( C/C \) also reflects some anisotropic motion. According to Dzuba (1996), the change in 2A\(_{zz}\) can be described by a librational model. Unlike rotational motion, in which the spin probe rotates entirely randomly, librational motion assumes that the spin probe rotates within a cone given by a small angle (\( \alpha \)). Therefore, it could be that the anisotropic motion seen in the \( t_R \) obtained from the \( C/C \) ratio arises from librational motion. Whether the change in 2A\(_{zz}\) with temperature is due to librational motion is currently under investigation. As ST-EPR is a laborious technique, more investigations are needed to make full use of this technique as a method to determine molecular mobility. Meanwhile, we will consider the measurements of 2A\(_{zz}\) as an estimate of molecular mobility.

**Molecular Motion and Intracellular Glasses**

As shown in Figure 4, a sharp increase in the molecular motion of CP was noticeable when the temperature of the sample was increased. Because the temperature corresponding to this sharp increase depended on water content, an attempt was made to explain this behavior according to the glass theory. From the plot of 2A\(_{zz}\) against temperature we derived two characteristic temperature points (see inset in Fig. 7 for details): at the intercept (midpoint \( T_g \)) and at the point of deviation from a straight line (onset \( T_{g'} \)). Figure 7 shows plots of the temperature at which the breaks occurred compared with the water content in pea axes. For cat tail pollen a similar type of plot was obtained (data not shown). At low water contents (<0.002 g water g\(^{-1}\) dry weight) the characteristic temperature points were difficult to determine exactly, as the legs below and above the glass transition did not show a sharp drop in 2A\(_{zz}\). The curves in Figure 7 are remarkably similar to state diagrams of intracellular glasses in seeds (Leopold et al., 1994; Leprince and Walters-Vertucci, 1995) and pollen (Buitink et al., 1996).

**Figure 5.** ST-EPR spectra of CP in pea axes at 0.08 g water g\(^{-1}\) dry weight recorded at various temperatures. Scans were recorded at 100 mW, a modulation frequency of 50 kHz, and a modulation amplitude of 5 G.

**Figure 6.** \( t_R \) of CP in pea axes at 0.08 g water g\(^{-1}\) dry weight as a function of temperature. The \( t_R \) values were obtained by comparing the \( L''/L' \) ratio and the \( C/C \) ratio with those of CP in anhydrous glycerol.
To ascertain whether the sharp changes in molecular motion were due to the melting of intracellular glasses, the onset $T_g$ was measured by differential scanning calorimetry and compared with the onset $T_g$ as determined by EPR (Fig. 8). Both curves closely matched one another, the EPR data being slightly lower than the differential scanning calorimetry data. The $T_g$ measured by differential scanning calorimetry has been found to occur above the $T_g$ measured at a molecular level (Kalichevsky et al., 1992). The midpoint of $T_g$ from EPR measurements was situated slightly above the midpoint of the $T_g$ measured by differential scanning calorimetry (data not shown). The $T_g$ for dry pollen (62°C) was lower than that for dry pea axes (92°C), which may be related to the high level of oligosaccharides in the latter (Amuti and Pollard, 1977; Saleki-Gerhardt and Zografi, 1994). The constant value of $T_g$ measured by differential scanning calorimetry when the last 2% of water was removed may indicate that the first small amount of water does not contribute to plasticization of the glass. It is possible that this water is not present in the glass but, rather, is located in some other part of the tissue (e.g. cell walls).

The change in molecular mobility during melting of intracellular glasses can be measured as a function of temperature or water content, the relationship being reflected by the state diagram (Fig. 8). To determine the change in molecular mobility as a function of water content instead of temperature, it is necessary to correct for the polarity change of the environment in which CP is present for each water content (Fig. 3). Therefore, using curves similar to those shown in Figure 4, the mobility at a certain water content was expressed as the difference between the maximum $2A_{zz}$ (at $-150^\circ$C, where the spin probe is assumed to be immobilized) and the $2A_{zz}$ measured at the desired temperature. We refer to this parameter as $\Delta A_{zz}$. Thus, an increase in $\Delta A_{zz}$ represents a relative increase in molecular mobility compared with the completely immobilized situation at $-150^\circ$C (i.e. the more the value departs from zero, the higher the molecular mobility). Figures 9 and 10 show the dependence of the molecular mobility ($\Delta A_{zz}$) on water content in pea axes and pollen, respectively. Between approximately 0.2 and 0.1 g water g$^{-1}$ dry weight, the mobility decreased with decreasing water content at all temperatures analyzed. When the tissues reached approximately 0.1 to 0.05 g water g$^{-1}$ dry weight, the $\Delta A_{zz}$ reached a constant value, indicating that the mobility had reached a minimum. When water contents were decreased further, below approximately 0.05 g water g$^{-1}$ dry weight, mobility increased again for pollen (Fig. 10). For pea axes, the mobility slightly increased again or reached a constant level at very low water content (Fig. 9). The water content corresponding to the minimum mobility (lowest $\Delta A_{zz}$) shifted to higher values with decreasing temperatures.
Behavior of Nitroxide Spin Probes in Dehydrating Organisms

To characterize molecular motion using EPR, a spin probe must be introduced into the material. Depending on the polarity of the spin probe, it will partition into the apolar oil phase, polar aqueous cytoplasm, or both. Several amphiphatic spin probes, such as TEMPO and 4-oxo-2,2,6,6-tetramethyl-1-piperidinyloxy, completely partition into the lipid phase during drying of cattail pollen (Hoekstra and Golovina, 1998). We found that the more polar 4-hydroxy-2,2,6,6-tetramethylpiperidinyloxy partially partitioned into the lipid phase of cattail pollen and pea axes during drying (data not shown). Because we were interested in the molecular mobility in the cytoplasm rather than in the lipid phase, we avoided the use of these spin labels and instead used the polar nitroxide spin probe CP.

There are several reasons why we believe that CP was not present in the lipid phase in the dry organisms. Upon melting of the oil, which occurred in cattail pollen at approximately −20°C and in pea axes at −40°C and −20°C, as determined by differential scanning calorimetry (data not shown), we never observed an abrupt appearance of a mobile EPR spectrum. Furthermore, during drying of pollen at 25°C, when the oil is liquid (Fig. 1), we did not observe the typical lipid signal with a hyperfine splitting constant of 14 G (Knowles et al., 1976). The resolution of the spectrum was high enough to distinguish a possible oil contribution from the aqueous contribution. When samples of elevated water content were heated above 80°C, two peaks were observed in the high-field part of the spectrum, one attributable to the water signal and the other to the lipid signal (data not shown).

During drying the hyperfine splitting constant of CP decreased from 16.5 to 15.5 G (Fig. 1). A decrease in hyperfine splitting constant is associated with a decrease in the polarity of the spin probe environment (Knowles et al., 1976). During drying the apparent polarity of the cytoplasmic environment decreased, possibly because of the decrease of the dielectric constant of the environment associated with the loss of water. This is also substantiated by the observation of the decrease in 2Δzz with water loss at −150°C (Fig. 3). It is interesting to note that cattail pollen and pea axes differ in the extent of their 2Δzz decrease.
Therefore, it seems evident that changes in water content are not solely responsible for this difference and that some intrinsic factors (e.g., salts or proteins) also contribute to the decrease in $2\Delta_{\text{zz}}$ during drying.

**Molecular Mobility in Biological Glasses**

Although the presence of intracellular glasses in seeds and pollen has been established, little is known about their viscosity and molecular mobility upon melting. We investigated the molecular motion of a spin probe in the cytoplasm of dry tissues to establish the relationship between glasses, molecular mobility, and storage stability.

During drying of cattail pollen and pea axes, the $\tau_R$ of CP increases from $10^{-11}$ s in the hydrated state to $10^{-2}$ s in the dry state. These values are consistent with values obtained in organic and inorganic glasses (Ito, 1983; Dzuba et al., 1984; Roozen et al., 1991), but contradict a previous study on molecular mobility in dry soybean axes, in which the $\tau_R$ of TEMPO was found to be approximately $10^{-10}$ s (Bruni and Leopold, 1990). Referring to the tendency of this spin probe to partition into oil during drying, we attribute this contradiction to the fact that these authors might have observed TEMPO in oillbodies. They nevertheless observed an abrupt change in molecular mobility around $T_g$. It is interesting that TEMPO in the lipid phase seems to be able to “sense” the glass transition of the cytoplasmic surroundings.

Molecular mobility was assessed in relation to water content and temperature. The change in distance between the two outer extrema of the powder spectrum ($2\Delta_{\text{zz}}$) with increasing temperature revealed a sharp increase in molecular mobility at a certain temperature that depended on the sample water content. This sharp increase was closely associated with the $T_g$ as measured by differential scanning calorimetry. When $\tau_R$ was monitored during melting of the glass, there was a decrease of four orders of magnitude, from $10^{-2}$ to $10^{-6}$ s.

We found a close correlation between changes in $\tau_R$, determined by the $C'/C$ ratio of ST-EPR spectra and changes in $2\Delta_{\text{zz}}$ derived from conventional EPR spectra during melting of intracellular glasses (compare Figs. 4 and 6). This correlation indicates that a change in $2\Delta_{\text{zz}}$ represents a change in molecular mobility. One can speculate on the type of molecular mobility that gives rise to the change in $2\Delta_{\text{zz}}$. Echo-detected EPR spectroscopy of nitroxide spin probes dissolved in organic glasses has revealed that the nitroxides undergo librational motions (Dzuba, 1996). This type of motion is described by a model that assumes that the spin probe rotates within a cone given by a small angle, $\alpha$ (Dzuba et al., 1992; Dzuba, 1996). Since we measured a $\tau_R$ for CP in intracellular glasses of $10^{-2}$ to $10^{-4}$ s, it seems unlikely that the change in $2\Delta_{\text{zz}}$ was due to a change in overall rotational motion; most likely it represents a change in librational motion, as has been established in wheat embryos (Dzuba et al., 1996). This is further supported by our ST-EPR study, in which some indication for anisotropic motion arising from libration comes from the different values of $\tau_R$ deduced from the line-height ratios $L'/L'$ and $C'/C$ (Fig. 6).

**Storage Stability in Relation to Molecular Mobility and Intracellular Glasses**

It has been shown that upon formation of glasses, the storage stability of seeds and pollen improves (Sun and Leopold, 1994; Sun, 1997; Buitink et al., 1998). The impact of intracellular glasses on the storage behavior of seeds and pollen has been ascribed to the high viscosity in the glass. Indeed, glasses are known to slow down detrimental reactions such as the rate of browning reactions (Karmas et al., 1992) and to increase the stability of enzymes (Chang et al., 1996). Although the presence of glasses has been associated with increased storage stability of seeds and pollen, there is not much known about the relationship between molecular mobility and storage stability. Our data on molecular motion in pea axes and cattail pollen in relation to water content and temperature enable a comparison with storage behavior, which is also known to depend on water content and temperature (Vertucci et al., 1994; Buitink et al., 1998).

We found a close relationship between the molecular mobility of CP and storage behavior in both pollen and...
seed axes. With decreasing water content molecular mobility, expressed as a change in the outer extrema of the EPR spectra ($\Delta A_{zz}$), decreased, whereas storage stability increased. Figure 11 clearly demonstrates that molecular mobility and storage stability are linked; the water content for optimum storage at various temperatures corresponds closely to the water content at which minimum mobility is observed (data from Vertucci et al., 1994; Buitink et al., 1998). Although the curves do not converge exactly with the 1:1 line, the slight deviation might be explained from errors in the determination of water content or the lack of an exact determination of the water content for optimum storage stability. Furthermore, the water content of minimum mobility was determined from the minima of the third-order polynomial equations. Especially at lower temperatures, the exact water content of minimum mobility is difficult to assess; there seems to be a plateau of minimum mobility present. To present the data clearly, we calculated a single value of minimum mobility, which should be considered with caution.

At low water content, molecular mobility seems to increase again. A similar observation has been made by Seitz et al. (1981), who observed that in Artemia cysts at lower hydration levels, the water self-diffusion coefficients increased slightly. Clegg et al. (1982) suggested from NMR studies on Artemia cysts that the increased mobility of water at a low water content might be due to a displacement of the water from polar-binding sites in the cell by sugars. Another explanation is that at these low water contents the spin probe partitions into a more mobile environment. However, it should be noted that this environment cannot be the lipid phase. CP in these phases will rotate faster than $10^{-8}$ s at room temperature. Therefore, the spectrum of CP in the oil phase will show the characteristic three sharp lines of a mobile spectrum and does not contribute to the $2A_{zz}$ of the powder spectrum.

However, other seed species show a divergence of the optimal storage stability from the $T_g$ curve (Sun, 1997). This is also true for cattail pollen (Buitink et al., 1998), in which we found that the optimum storage conditions coincided with the water content at which molecular mobility was minimum (Fig. 11). This minimum mobility occurred below $T_g$. Therefore, we propose that measurements of molecular mobility rather than state diagrams be used to predict optimum storage conditions.

Based on the relationship between the minimum molecular mobility and the water content of optimal storage, we made an attempt to predict the optimum storage conditions at subzero temperatures. This was found to be $-20^\circ$C and $-60^\circ$C for pea axes (Fig. 9, E and F) and $-20^\circ$C and $-40^\circ$C for cattail pollen (Fig. 10, E and F). In practice it will not be possible to analyze storage behavior at these low temperatures on a realistic time scale. At $-20^\circ$C the minimum molecular motion of pea can be estimated at approximately 0.10 g water g$^{-1}$ dry weight, and at $-60^\circ$C between 0.14 and 0.2 g water g$^{-1}$ dry weight. A similarly elevated optimum water content for low-temperature storage was predicted by Vertucci and Roos (1993) on the basis of thermodynamic considerations. At $-20^\circ$C the minimal molecular motion of pollen was approximated at 0.1 g water g$^{-1}$ dry weight, and at $-40^\circ$C between 0.15 and 0.2 g water g$^{-1}$ dry weight. With lower temperatures, an increase in the water content at which minimum mobility was observed became evident. This implies that too much drying increases mobility and reduces longevity and should be avoided, particularly when cryogenic storage is considered. Where determinations of optimal storage conditions of seeds and pollen by germination assays take too long to perform, measurements of $2A_{zz}$ might be considered instead.

**Optimum Storage Conditions Predicted by Molecular Mobility**

Recent studies report a water-content limit below which seed longevity did not increase further (Ellis et al., 1989, 1990), and which had an adverse effect on seed viability and seed vigor (Vertucci and Roos, 1993; Vertucci et al., 1994; Buitink et al., 1998). It is thought that the removal of the last remaining water molecules may destabilize biological structures (Sun, 1997; Buitink et al., 1998) or enhance the lifetime of free radicals due to the loss of water as a quencher (Karel, 1975). However, we observed an increase in molecular mobility for cattail pollen (which also occurred to a lesser extent in pea axes) when it was dried to very low water contents. This increase in molecular mobility also might be responsible for the decreased storage stability observed at these low water contents.

In some cases, state diagrams can be used to predict the optimal storage conditions (Sun, 1997). For pea axes the optimum water contents of storage at a certain temperature were found to coincide with $T_g$ (Fig. 8; Vertucci et al., 1994).

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