Mechanical Properties of Plant Cell Walls Probed by Relaxation Spectra

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Transformants and mutants with altered cell wall composition are expected to display a biomechanical phenotype due to the structural role of the cell wall. It is often quite difficult, however, to distinguish the mechanical behavior of a mutant’s or transformant’s cell walls from that of the wild type. This may be due to the plant’s ability to compensate for the wall modification or because the biophysical method that is often employed, determination of simple elastic modulus and breakstrength, lacks the resolving power necessary for detecting subtle mechanical phenotypes. Here, we apply a method, determination of relaxation spectra, which probes, and can separate, the viscoelastic properties of different cell wall components (i.e. those properties that depend on the elastic behavior of load-bearing wall polymers combined with viscous interactions between them). A computer program, BayesRelax, that deduces relaxation spectra from appropriate rheological measurements is presented and made accessible through a Web interface. BayesRelax models the cell wall as a continuum of relaxing elements, and the ability of the method to resolve small differences in cell wall mechanical properties is demonstrated using tuber tissue from wild-type and transgenic potatoes (Solanum tuberosum) that differ in rhamnogalacturonan I side chain structure.

The cell wall of higher plants is a composite material consisting of threadlike, partially crystalline cellulose microfibrils embedded in a hydrated matrix of pectins, hemicelluloses, and glycoproteins. In vascular plants, more than 200 genes encoding glycosyl transferases, and an unknown number of genes from other families, are probably devoted to the synthesis and assembly of cell wall polysaccharides. The structural complexity of plant cell walls may seem surprising, if conveying mechanical strength to the plant body were their only role, but this is not their only role, as displayed in the review by Jarvis and McCann (2000).

It has long been recognized that permitting the cell wall to expand during growth, and coordinating this expansion with concurrent deposition of new wall material, requires rather sophisticated control over wall rheological properties. “Properties” here refer to diverse phenomena, not only purely physical responses to stresses but also mechanical behavior that arises from enzymatic processes involved in growth and that may be lost in isolated or “dead” wall preparations. Cosgrove (1993) reviewed methods that have been used to gauge these respective aspects of cell wall mechanics, and Schopfer (2006) discussed pitfalls in data interpretation that can result from not appreciating this distinction. Yet, ever since Kamiya et al. (1963), it has been clear that plant cell walls exhibit viscoelastic properties like retarded elasticity, and most workers since Probine and Preston (1962) have assumed that these properties should be important to cell growth and/or other aspects of plant cell function that depend on cell wall behavior (Thompson, 2008).

The discoveries of xyloglucan endotransglycosylases (Nishitani and Tominaga, 1992; Fry et al., 1992) and expansins (McQueen-Mason et al., 1992) stimulated research into the biochemical aspects of cell enlargement and a search for gene products that stimulate it, while characterization of cell wall material properties concurrently tended to receive less attention. Cosgrove (1993) foresaw, however, that the ability to alter single wall components in controlled ways (e.g. using mutants or transformants) would allow us to relate wall structure and polymer composition to both rheological...
theory and physiological function. Mechanically significant structural changes in mutants may not necessarily be easily detectable by determining monosaccharide profiles (Bosca et al., 2006) or from other types of relatively simple compositional analysis. Architectural modifications may occur that impact wall mechanical properties and are detectable by appropriate biophysical methods but for which all but the most detailed chemical analyses fall short. Biophysical analysis has yet to be employed as a screen to identify cell wall mutants, but a number of characterized mutants have been found to have altered cell wall mechanical properties (Reiter et al., 1993; Ryden et al., 2003; Peña et al., 2004; Zhong et al., 2005).

Although mechanical measurements on cell walls, stimulated by the issue of how walls extend during growth, go back at least to the Dutch botanist A.N.J. Heyn (1933), in most of this work, load or stress was recorded at a fixed rate of extension (Cleland, 1967), which does not afford a clear measure of time-dependent rheological properties. Measurement of “creep” under a constant load (Probine and Preston, 1962) can reveal these, but probably the most incisive type of biomechanical analysis, and the one most capable of detecting subtle changes in cell wall rheology, is to determine stress relaxation spectra.

Stress relaxation is a time-dependent decline in stress (or load) when a material is held at a constant, initially load-induced, mechanical deformation or strain. Stress relaxation, and retarded elasticity, reflect viscosity-retarded changes in conformation or position of load-bearing polymer chains. Different changes of this type can occur over widely different time scales, depending on how local or long range these chain displacements are and how large a viscosity consequently restricts them. Mathematical analysis of overall behavior is achieved by considering the behavior of analogous mechanical models containing multiple elements, as in Figure 1, A (for stress relaxation) and C (for retarded elasticity). Any given type of viscosity-restricted conformational change (a single element in Fig. 1, A or C) possesses a characteristic relaxation or retardation time (τ), or time to proceed exponentially all but [1/e]th, or 63.2%, of the way to completion. A relaxation spectrum displays the distribution of different τ values (and hence of rheologically differing mechanical elements) in the material and their relative importance. The “mechanical elements” are often modeled by arrangements of springs and “dashpots,” as in Figure 1. Figure 1A represents the model behind BayesRelax, that is, where the wall polymers are represented by a continuous spectrum of relaxing components (approximated by 100 elements in the algorithm). To deduce a relaxation spectrum from stress relaxation measurements in the most general way, one starts with the possibility that the material could possess many different relaxation mechanisms with τs extending over a wide range; the peaks, at particular τ values, in the spectrum reflect which of these mechanisms are mechanically more important. A specific wall polymer modification that affects just one particular relaxation mechanism might thus be expected to alter some particular part of the relaxation spectrum, even if it does not greatly change the mechanical behavior of the wall overall. This is what makes stress relaxation spectral measurements potentially more sensitive, for detecting subtle changes in cell wall mechanical properties, than simple rheological measurements are.

The earliest measurements of stress relaxation in plant cell walls of which we are aware were made by Haughton et al. (1968) and Haughton and Sellen (1969) on four species of algae. Shortly thereafter, Cleland and Haughton (1971) reported stress relaxation curves for oat (Avena sativa) coleoptile cell wall skeletons (intact wall structure of killed tissue). Throughout that decade, Yamamoto, Masuda, and coworkers pursued stress relaxation measurements with coleoptiles and other higher plant cell walls (Yamamoto et al., 1970; Yamamoto and Masuda, 1971; Fujihara et al., 1978; Sakurai et al., 1982), deducing relaxation spectra from simple time courses of stress relaxation. In hindsight, this was a relatively insensitive basis for obtaining relaxation spectra, and these workers lacked more recently developed mathematical tools for analyzing stress relaxation.

A more incisive method for deducing relaxation spectra, applicable to a wider range of relaxation times than can be covered by simple relaxation time courses, is “dynamic” measurement of the variation of stress under an oscillating (usually sinusoidally varying)
strain. Dynamic measurements at a single frequency, which have widely been used on food materials and wood and in a few instances on thin-walled plant tissues (Ramana and Taylor, 1994; Takeda et al., 2002), are interesting but do not enable a relaxation spectrum to be deduced. This is possible only if dynamic measurements are made over a range of oscillation frequencies (Findley et al., 1976), which is often called a “frequency sweep.” The main advantages of this method are that (1) rapid relaxation processes (having subsecond relaxation times) can be detected and quantified, and (2) relaxations with longer relaxation times can be characterized more accurately than when they are deduced from small deviations of a time course from a simple exponential decline of stress. The method has been used for analyzing the mechanical properties of wood (Ouis, 2002, and refs. therein) and is introduced for thin, primary plant cell walls in this work. (Frequency sweep measurements were made previously by Whitney et al. [1999] on homogenized tomato [Solanum lycopersicum] fruit cell walls and bacterial cellulose/xyloglucan composites, but they did not deduce relaxation spectra from their data.)

Determining relaxation spectra from frequency sweep biophysical data is not trivial; it is what is termed an “ill-posed problem,” meaning that experimental error due to imprecision of measurement may permit many, including some quite different, solutions. Methods to handle ill-posed problems have been developed (Tikhonov and Arsenin, 1977) and are frequently used in the physical sciences, but apart from limited use in food science, they have thus far received rather little attention in biology.

In most applications, an ill-posed relaxation spectrum problem is “solved” by imposing upon the connection between the data and the spectrum an additional constraint called “regularization.” For example, the “smoothness” regularization constraint gives preference to smoother solutions, suppressing the emergence of spurious spectral peaks. We take this approach here, applying the most recent advances in handling ill-posed problems (Hansen, 2008) to the analysis of cell wall rheology in intact plant tissue. The only spectral peaks allowed by smoothness regularization are those that are required to obtain a satisfactory fit to the experimental data. This conservative approach means that data sets with small relative variance will usually give more feature-rich relaxation spectra than more noisy data sets will. The computer program developed for this study, BayesRelax, as well as a user’s guide, are made available to the scientific community through the Web site at www.BayesRelax.org.

We employ potato (Solanum tuberosum) tuber tissue, both wild type and transgenic tissue with altered rhamnogalacturonan I (RG-I) side chains, as a case study of the use of BayesRelax. Compared with the wild type, the transformants are reduced either in RG-I β-1,4-galactans (line T13.1; Sørensen et al., 2000) or in RG-I α-1,5-arabinans (line T7; Skjøt et al., 2002). Previous investigation (Ulvskov et al., 2005) by more conventional methods suggested that T13.1 cell walls differ mechanically from those of the wild type. Potato tubers afford ample amounts of relatively homogenous tissue suitable for frequency sweep measurements in standard rheometers used in the food industry. Our experience with this material suggested that its cell walls’ natural state of hydration and of normally imposed tension (due to turgor pressure, hereafter denoted P) are probably important factors in its mechanical behavior. Therefore, we consider that it is biologically relevant to make these measurements on living, normally hydrated and turgid, tissue. However, the use of such tissue involves potential complications, resulting both from the possibility of progressive metabolic modification of the cell wall during the measurements and from the dependence of a turgid tissue’s elastic properties upon P, which could change for any of several reasons during the measurement. Problems raised by changes in P are considered in both “Results” and “Discussion” and are dealt with in detail in Supplemental Data S1. We demonstrate the successful determination of relaxation spectra that distinguish the transformants from the wild type, despite data that are rather noisy, apparently due primarily to the P problem.

RESULTS

Dynamic Stress Relaxation Measurements

Operation of the rheometer yields, for each strain-oscillation frequency (ω), a value for what is called the complex modulus and its two components, storage modulus and loss modulus. Storage modulus (G'') reflects that part of the deformation work that can be recovered and hence represents the elastic component...
of the mechanical properties, while loss modulus ($G''$) reflects the fraction of the deformation work that cannot be recovered but is dissipated as heat, due to viscous flow in the material. A relaxation spectrum can be validly derived from these data only if $G'$ and $G''$ do not vary as a function of small changes in deformation (strain). Therefore, a strain sweep (varying the magnitude of the oscillatory imposed strain) was performed to determine if potato tissue possesses a linear elastic range. $G'$ and $G''$ were nearly constant up to a strain of 0.1%, the limit of linear viscoelasticity (Fig. 2). The rheometer was subsequently operated in constant maximum stress mode with maximum shear stress preset to 26 Pa. This setting led to maximum shear strains in the range of 1 to $5 \times 10^{-4}$.

Frequency sweep measurements were then performed using these settings. Figure 3 (A–C) shows the mean $G'$ and $G''$ values obtained from all the employed frequencies, plotted against $\omega$ on a log scale, for the wild type and the two transformed lines with modified cell walls. The curve that is drawn for each data set was fitted to the data as described in the next section. As shown by the error bars, variance among replicate measurements at particular frequencies was often quite large. Great variability was similarly noted by Alvarez and Canet (2000) in creep time courses on potato tissue under shear stress, obtained using a comparable rheometer.

An important source of variance in our data is related to the variable normal force (NF) to which the tissue specimen is subjected in order to hold it firmly between the oscillating plates of the rheometer during the measurement. Although the rheometer was set to deliver a target NF of 30 g, the NF that the instrument initially reported, which was often higher (and sometimes much higher) than 30 g, subsequently relaxed, rapidly during the 1-min premeasurement equilibration period and usually for approximately the first 100 s of measurement, then more gradually. Since the force that a turgid plant cell exerts against a compressive strain imposed externally against its cell wall varies with $P$ (Davies et al., 1998; Lintilhac et al., 2000; Wei et al., 2001), the decline in NF very likely reflects a decline in $P$. Variations in $P$ are the only reasonable, identifiable source for the large variations in modulus values that the instrument reported, since plant tissue rigidity or elastic modulus varies with $P$ over a very wide range (Falk et al., 1958; Niklas, 1988; Davies et al., 1998; Wei et al., 2001). This is apparently why the modulus values at any strain-oscillation frequency correlate strongly with NF. Figure 4 gives two examples of this correlation, and its inset shows the slopes and correlation coefficients ($r^2$ values) for similar plots of $G'$ versus NF for all the applied $\omega$ values (here, and in the following, data from originally applied $\omega$ values that differed by less than 10% have been

![Figure 3](image-url). Frequency sweep measurements in the range 0.001 to 10 Hz for potato tuber discs of dimensions 15 × 3 mm. Error bars indicate SD. Solid lines represent fitting of storage modulus ($G'$) and dashed lines represent fitting of loss modulus ($G''$) by Fredholm equations with parameters deduced as explained in the text. A to C show raw data, and D to F show the data after normalizing to an NF of 30 g, as described in “Materials and Methods.” A and D, The wild type. B and E, T7.2. C and F, T13.1.
Figure 4. Examples of the correlation between $G'$ and NF values at two particular frequencies within the range of the measurements. White symbols represent $3 \times 10^{-3}$ Hz, and black symbols represent 10 Hz. Different symbols show data from the different genotypes: the wild type (triangles), T7.2 (circles), and T13.1 (squares). Similar correlations were obtained at other frequencies, for $G''$ as well as $G'$. The inset table lists the slopes of the regressions and the correlation coefficients ($r^2$ values) for $G'$ for all the employed frequencies.

combined for further analysis; see “Materials and Methods”.

The variance in $G'$ and $G''$ values, therefore, was reduced by normalizing the data to a standard NF value of 30 g. Normalization was performed by linear regression separately for each of the nominal frequencies for the pooled data from all genotypes. Ignoring possible differences between genotypes avoids introducing, in the normalization, any bias between different genotypes that might either lead to, or increase, an apparently significant difference among their relaxation spectra. The data thus normalized, plotted in Figure 3 (D–F), show considerably reduced variance, as expected.

Relaxation Spectra

Relaxation spectra are typically derived from dynamic measurements using a generalized Maxwell model comprising a large (ideally an infinite) number of Maxwell elements in parallel (Ferry, 1980), as in Figure 1A. For solids, which presumably include non-growing cell walls, one of the parallel elements is simply a spring, which precludes any steady, irreversible flow. A relaxation time, $\tau$, is associated with each of the remaining elements $i$, with $\tau_i = \eta_i / E_i$, where $\eta_i$ is the viscous resistance of dashpot $i$ and $E_i$ is the tensile modulus of spring $i$. Each of these components contributes to relaxation over about two decades of log (time); the $\tau$ for any one of them is the time needed for that one to proceed $(1 - 1/e)^{th}$, or 63.2% of the way toward complete relaxation. The presently used rheometer imposes shear strain on the material rather than extension, so we deal with shear moduli ($G$) rather than the tensile moduli ($E$) that would nominally apply to the model in Figure 1A, but the mathematics of the relationship between the relaxation spectrum and the modulus values are just the same.

A relaxation spectrum $H(\tau)$ depicts the relative contributions of Maxwell elements (Fig. 1A), with different $\tau$ values, to the overall relaxation capability of the tissue. $H(\tau)$ does this, for any given $\tau$ value $\tau_v$ by representing this contribution as the increase in overall modulus ($dG$) per infinitesimal bit of increase in log $\tau$ in the neighborhood of $\tau_v$ that is, $(dG/d \log \tau_v)$. Since $G$ has units of Pa and log $\tau$ is dimensionless, $H(\tau)$ has units of Pa.

As described in “Materials and Methods,” the relationship between a relaxation spectrum and the $G'(\omega)$ and $G''(\omega)$ values obtainable from frequency sweep data is given by a pair of equations called the Fredholm equations. For each of our data sets, the BayesRelax program derived a relaxation spectrum by obtaining, by a method of successive approximations, the $H(\tau)$ function that, when tested in the Fredholm equations by numerical integration, gave the best fit to the given set of $G'(\omega)$ and $G''(\omega)$ values, subject to the regularization constraint that was imposed. The $G'(\omega)$ and $G''(\omega)$ profiles that could be calculated from the Fredholm equations, using the thus-deduced $H(\tau)$ functions, are the curves drawn through the data points in Figure 3. In deriving each relaxation spectrum, the continuum of relaxing elements in the generalized Maxwell model was approximated with 100 discrete relaxation times.

Figure 5 shows the entirety of each spectrum as fitted to the raw data (Fig. 3, A–C) using the smoothness regularization constraint mentioned previously.

Figure 5. Relaxation spectra corresponding to data in Figure 3 (A–C) fitted using BayesRelax with the smoothing constraint. Squares indicate the wild type, white diamonds indicate T7.2, and black diamonds indicate T13.1. Bars show the 67% probability range for each calculated point (see “Materials and Methods”).
Figure 6 gives comparable spectra obtained from the data after these were normalized to the standard NF of 30 g (Fig. 3, D–F) as explained above. This normalization obviously did not change the basic form of the spectra, although it reduced the error bars associated with some portions of the curves and thereby increased somewhat the extent to which the three spectra appear to differ from one another significantly. Each error bar represents the range of $H(t)$ values within which the true value of $H(t)$ for that point falls with a probability of 67% (see “Materials and Methods”).

In Figures 5 and 6, only the range between $\tau = 0.1$ and 1,000 s, corresponding roughly to the range of frequencies used in our frequency sweep, can be regarded as reliably fitted. In the range between 0.1 and 1,000 s, the relaxation spectrum is mainly determined by the constraints from the measured data, while the form of the estimated plot outside this interval is heavily influenced by the smoothness constraint and by a requirement that $H(\tau)$ fall to 0 at the lower and upper limits that must be set for the spectrum (beyond the range of the actual data; see “Materials and Methods”) in order to perform the Fredholm equation integrations. Hence, the interval between 0.1 and 1,000 s is marked “reliable range” in Figure 5, and its limits are indicated with vertical dashed lines in Figure 6. Due to the correlations between neighboring points in the relaxation spectrum that is introduced by the smoothness constraint, the reliable range may actually extend slightly beyond the indicated interval. However, as the extent of valid extrapolation is somewhat unclear and depends on the noise level in the data, we indicate only the “traditional” reliable range in Figures 5 and 6.

The spectra resolve two peaks of relaxing components: a quickly relaxing one with $\tau$ values shorter than about 2 s (apparently peaking at approximately 0.1 s), and a slowly relaxing peak with $\tau$ values from 10 s up to approximately 200 s. The quickly relaxing peak behaved relatively similarly in all three potato lines. On the other hand, compared with the wild type, line T13.1 shows a significantly reduced relaxation intensity at $\tau$ values between 200 and 1,000 s in both Figures 5 and 6, plus an apparently significant increase (Fig. 6) between $\tau = 0.1$ and 0.6 s. The latter might represent a downshift (to lower $\tau$ values) of at least part of the relaxation capability that has disappeared from the 200- to 1,000-s region. Small but significant (judging from the error bars) differences between the wild type and T7.2 occur in two parts of the reliable range in Figures 5 and 6 and might be interpreted as a minor part of T7.2 relaxation capability having been shifted from the 10-s region down into the 0.75-s $\tau$ region.

The consequences of applying the smoothness constraint that was used for the spectra in Figures 5 and 6 can be illustrated by comparing the results given there with those of a different constraint that is often used, the maximum entropy method (Elster and Honerkamp, 1991). Figure 7 is equivalent to Figure 6, except for the fitting constraint that was employed. The maximum entropy constraint favors low peak height and, as a side effect, tends to favor solutions with extra peaks and shoulders. These features may reflect something real, but they may instead, if there is appreciable experimental error, be artifacts of fitting that lead data interpretation astray. The plots in Figure 7 agree in general with those in Figure 6, depicting line T13.1 as having a substantial drop in $H(\tau)$ below that of the wild type in the $\tau$ range above about 200 s. However, Figure 7 displays a shoulder of increase in T13.1 $H(\tau)$ above the wild type value around $\tau = 3$ s, which is found in neither Figure 5 nor Figure 6.
spectra of Figure 7 are shown without error bars for clarity and serve mainly to confirm, using a different method of analysis, the general features of the relaxation spectra of the three potato lines.

DISCUSSION

Our measurements of stress relaxation in potato tuber tissue by the “dynamic” (sinusoidally oscillating strain) method yield a prominently two-peaked relaxation spectrum (Figs. 5 and 6). Contrary to possible naive impression, the peaks do not represent times at which there is a maximum rate of relaxation. The rate of relaxation of all Maxwellian elements is actually maximum at the start of the process; the τ of any element is the time at which it will have gone 63.2% of the way to completion. H(τ) plotted on the ordinate is the rate of decrease in modulus (or in stress) relative to log(time); thus defined, for any given relaxing element, H reaches a maximum at the time (after strain and stress are imposed) equal to this element’s τ. A peak in H(τ) can be caused by simply a shoulder, or an inflection, in the change in modulus with time, or in the increase in moduli with ω. A peak in H in the log(τ) plot means just that more total relaxation occurs in that decade of log time than in the decades to either side. The H(τ) minimum at τ values between 1 and 10 s, which separates the two peaks in our spectra, is brought on by the minimum in G” and the slight inflections in the curve for G” in the ω range around 0.2 to 2 s⁻¹ in our frequency sweep data (Fig. 3).

A two-peaked relaxation spectrum is characteristic of amorphous, synthetic polymers with chains long enough to create “entanglement coupling.” This is where occasional kinking of long, randomly coiled chains around each other creates a transient network structure (or alternatively, where movement (“reptation”) of very long chains within the “sheath” of adjacent chains that confines them is retarded by an extralong resistance compared with that for shorter range molecular motions; Ferry, 1980). The peak at a low τ value reflects local movements of chain segments, while the large τ peak reflects a much slower slippage of entire chains past coupling points (or these chains’ reptation) under an imposed stress. In entanglement coupling, the respective spectral peaks, however, often seem to be separated by about five decades of log τ (Ferry, 1980), in contrast to our peaks, which are only about three decades apart.

Unlike an amorphous polymer, as noted in the introduction, cell walls involve structure, namely, cellulose microfibrils and intervening matrix polymers, which might instead be responsible for multiple relaxation spectral peaks. When a wall is strained elastically, its virtually inextensible microfibrils must become displaced relative to one another, either by separation or (more generally) by slip (lengthwise movement of a microfibril relative to an adjacent, more or less parallel, one). The former would require modest extension straining of the matrix, while the latter must locally shear strain, much more strongly than the macroscopic strain in the wall, the matrix intervening between adjacent, more or less parallel, fibrils.

Whitney and Cosgrove (1989) reported stress relaxation spectra for cell wall specimens from killed cucumber (Cucumis sativus) hypocotyls, which showed a

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broad, but not box-like, maximum centered between 0.2 and 0.6 s (in different spectra) and a minimum at near 100 s, the increase above which did not reach a maximum by the spectrum’s upper limit of 200 s but presumably would have given a second peak beyond this point. These spectra thus appear to be two peaked, like ours, but right shifted to higher \(\tau\) values (relatively slower rates of relaxation) than those of our peaks. This shift is to be expected from measurements on killed, turgorless tissue as against our turgid potato cells, because, as previously noted, \(P\) increases a tissue’s elastic moduli and \(\tau\) is inversely related to modulus.

More recent authors have used time courses of creep under a steadily applied load to deduce rheological properties of thin-walled plant tissues. Although the same mechanisms participate in creep as in stress relaxation, a feasible analysis of retardation uses not the generalized Maxwell model (Fig. 1A) but instead a “Burgers” model (Fig. 1C), which comprises a number of “Kelvin” or “Voigt” elements (Fig. 1B) in series along with one Maxwell element. The latter allows for instantaneous (unretarded) elasticity and for the possibility of steady flow (its dashpot’s viscosity being infinite if steady flow cannot occur), while the former represent multiple retarded-elastic straining mechanisms with different retardation times (times for 63.2% extension under a fixed load). The retardation time of any given structural element is typically longer than its relaxation time, because during relaxation, extension of any Kelvin/Voigt element’s spring is opposed by compression of other springs in series with it in a Burgers model, whereas during retarded extension, it is not so opposed, which allows it to approach equilibrium extension more gradually. Because of this and the fact that retarded elasticity is measured as compliance, which is the reciprocal of the moduli involved in relaxation, the relaxation and retardation spectra of a given material usually do not resemble each other closely but tend to have peaks and valleys at \(\tau\) values that are not remotely distant from one another (Ferry, 1980).

Alvarez and Canet (1998, 2000) and Thompson (2001, 2005, 2008) analyzed creep data using a Burgers model comprising two Kelvin elements in series with a steady-flow viscosity. The materials that Thompson tested are not structurally comparable with potato tissue, but Alvarez and Canet used living, turgid potato tissue equivalent to that studied here. From creep curves that extended over just 2 min, they inferred \(\tau\) values, in different measurements, ranging from about 100 to 700 s for one Kelvin element and 14 to 62 s for the second one (which was not consistently detected; Alvarez and Canet, 2000). Their higher \(\tau\) component falls within the general \(\tau\) range of the slower (longer \(\tau\)) peak in our relaxation spectra (Figs. 5 and 6), but their lower \(\tau\) component falls in the range in which our spectra display a minimum. As is evident from the mentioned numbers and from their statements (Alvarez et al., 1998; Alvarez and Canet, 2000), the results of their curve fitting for any given material were extremely variable and thus presumably inaccurate. Also, the one actual creep time course that they published appears to lack the initial part of the post-loading response, which would be needed to detect any retarded-elastic straining having \(\tau\)s in the range less than 1 s, comparable to the lower \(\tau\) peak in our spectra.

We consider that simple creep as well as simple stress relaxation time courses for cell walls, even if low in noise, are not data rich enough to distinguish specific rheological models, like those of the foregoing authors, from a generalized Burgers or Maxwell model involving a spectrum of retardation or relaxation times, as has generally been found necessary for polymeric materials (Ferry, 1980). To obtain data that are rich enough in detail to adequately define such spectra, or to justify models with only a limited number of components, it seems necessary to employ dynamic (frequency sweep) stress relaxation measurements and to analyze them assuming a near continuum of possible relaxation times.

**Problems in Stress Relaxation Measurements**

Without a regularization procedure, fitting experimental data to a model containing more parameters than the amount of information in the data will lead to instability in the fitting routine, giving parameter values that cannot be trusted. An apparently good fit can then be due merely to the extra parameters employed. For a reliable analysis, the number of fitting parameters should not exceed the effective number of degrees of freedom in the data. This has often been overlooked in stress relaxation work.

The emulsified or homogenized preparations from which relaxation spectra of synthetic polymers are commonly obtained (Tan et al., 2000) have been imitated for plant tissue by homogenizing, ethanol washing, drying, and rehydrating its cell walls (Kunzek et al., 1997; Whitney et al., 1999). However, such preparations are probably not very relevant to the biological properties of the cell walls of living tissue. For one thing, completely dried cell walls most likely cannot be fully reconstituted, by rehydration, back into their native rheological condition. The analogous problem is termed “retrogradation” in the chemistry of solubilized polysaccharides and “hornification” in wood pulp fiber technology (Fernandez Diniz et al., 2004).

We have instead shouldered the complications of using living tissue in order to study cell walls in their native states of hydration and physical condition. These complications comprise, first, biochemical activities of the tissue, including the possibility of irreversible cell wall extension (growth) as well as wound reactions to cutting of tissue discs, and second, problems with turgor pressure and osmotic relaxation.

The postharvest metabolism of RG-I side chains in potato tubers (Bush et al., 2001) implies that the walls
might not remain in their initial biochemical state during a protracted incubation of isolated potato discs during stress relaxation measurement. We guarded against this by keeping pretreatment and measurement periods as short as practicable (1 min and 20 min or less, respectively). The range of $\omega$ values used here were limited, on the slow end, by this short time requirement. On the fast end, usable $\omega$ values were limited by an apparent tendency of the tissue discs to begin slipping between the rheometer plates (as indicated by $G'$ values falling with $\omega$) at $\omega$ values greater than 10 s$^{-1}$, despite the rheometer’s arrangements to hold specimens firmly, whose consequences are noted below.

Walls of turgid cells may differ mechanically from those of nonturgid cells in that their walls are compressed in the normal (perpendicular to the cell surface) direction by $P$, potentially affording more and/or closer interpolymer contacts within the wall structure than would occur in fully relaxed walls. This would probably increase the effective viscosities governing stress relaxation and could also affect their degree of hydration, which is probably also important to these viscosities. Thus, it is biologically relevant to try to probe walls of turgid cells, as was done here. However, most of the problems encountered in interpreting dynamic rheological measurements on a living plant tissue are related to its $P$. This is because a plant tissue’s rigidity or elastic modulus varies strongly with its $P$, as noted above, and $P$ can change during the measurements. Because most of these problems are relevant not only to this work but would need to be faced in any future frequency sweep measurements on living plant tissues, it is desirable to consider them. However, due to their number and physiological complexity, to analyze them adequately requires considerably more space than can be devoted to it here. Therefore, we undertake this in Supplemental Data S1. Table I here provides a key to the topics considered there and gives, in the right column, our conclusions, from that analysis, as to which aspect(s) of this work each listed problem actually affects.

### Rheological Changes in Tissues with Modified Wall Polymers

The regularization method employed to obtain Figures 5 and 6 (which we recommend) is very conservative with regard to disclosing differences. However, it is able to differentiate all three potato genetic lines. Potato line T13.1 is reduced by 70% in RG-I galactans, which translates into a 6% overall change in cell wall composition compared with the wild-type line Posmo (Sørensen et al., 2000). We previously inferred that in T13.1, a slowly relaxing component had been lost (Ulvskov et al., 2005). Our relaxation spectra here confirm this, indicating that in T13.1, a component with a $\tau$ value of around 1,000 s is greatly reduced. However, according to Figure 6, components with $\tau$ values in the 6- to 30-s and the 0.2- to 0.5-s ranges have increased in total by about the same amount as the decrease in the $\tau$ approximately 1,000-s region.

$^{13}$C-NMR measurements have been interpreted as indicating that galactans are among the most freely mobile polymers in hydrated, pectin-rich primary walls of onion bulbs and a few other plant materials (Foster et al., 1996; Ha et al., 1997), including potato (Tang et al., 1999). Thus, pectic galactans might be expected to have short relaxation times, rather than $\tau$ values near the upper limit of our spectrum. The $\tau$

### Table I. Physiological/technical problems for dynamic rheological measurements on turgid tissues

These problems are analyzed in Supplemental Data S1.

| No. | Problem | Potential | Actual |
|-----|---------|-----------|--------|
| 1   | “In vivo stress relaxation” related to a cell’s capacity for irreversible wall expansion (cell growth)$^b$ | A, B | (B)$^c$ |
| 2   | Osmotic relaxation of $P$ after a change in $P$ imposed by rheometer action | A, B | $0^d$ |
| 3   | Change in $P$ due to one or more of the viscoelastic mechanisms of wall stress relaxation$^e$ | B | $0^d$ |
| 4   | Compression, and resulting collapse, of cells impacted by the rheometer sample grip or antislip arrangements$^f$ | B | B |
| 5   | Leakage, from cells that collapse due to No. 4, of solutes into a tissue’s cell wall space, causing water loss from other cells | B | B |
| 6   | Evaporative water loss from a tissue sample’s edges having unprotected contact with ambient air | B | B |
| 7   | Variations in thickness of tissue discs$^g$ and in pressure applied to them by sample gripping arrangements$^h$ | C | C |

$^a$Pertinence of listed problem to any of the following: A, direct contribution to relaxation spectrum; B, $P$ relaxation during measurement run, leading to decline in NF and resulting scatter of $G'$ and $G''$ values at a given $\omega$; C, effect on NF that should not correlate consistently with $G'$ and $G''$, contributing to scatter of NF-normalized $G$ values. $^b$Cosgrove (1985, 1987). $^c$Only a marginal influence, at most. $^d$Potential problem that we conclude (in Supplemental Data S1) does not influence our measurements on potato discs but could affect other work. $^e$Potential problem for potato relaxation spectrum because NF rapidly declines over the same time scale as the spectrum’s longer $\tau$ peak. $^f$In this work, local compression of cells by antislip knurls on the rheometer’s pressure plates. $^g$Variations in disc thickness cause variations in initial NF if pressure plates cease advancing at exactly the target separation. $^h$Pressure plate behavior in the rheometer used here is not under the operator’s manual control and seems variable, contributing to variations in initial NF.
approximately 1,000-s component that is reduced in T13.1 thus probably reflects an indirect effect of the missing galactan on other wall components.

Galactan side chains, although themselves mobile, might restrict the RG-I polymer backbone mobility, as branches or side chains on synthetic polymers typically do (Ferry, 1980). In that case, shortening or eliminating many of these side chains might reduce the viscous resistance that retards RG-I backbone motion. Since the larger the value of $\tau$, the higher the associated viscous resistance, this change would downshift the part of the spectrum that reflects RG-I backbone mobility, corresponding to the apparent downshift noted in “Results.”

On the other hand, RG-I side chains might act as hydrated spacers within the wall, limiting direct contacts between RG-I and homogalacturonan backbones and possibly also between cellulose microfibrils. Eliminating side chains would then increase these associations and stiffen the wall, as indicated for arabinan side chains by Jones et al. (2003) for stomatal guard cell walls, and as our previous experiments suggested (Ulvskov et al., 2005). Wild-type component(s) with $\tau$ values approximately 1,000 s would, in T13.1, have become immobile enough that their relaxation lies beyond the upper $\tau$ limit of our spectrum’s reliable range. This is suggested by the sharp rise in $H$ with $\tau$ in the 5,000- to 10,000-s range of the T13.1 spectrum (Figs. 5 and 6), although since this is above the reliable range, this feature is not compelling. In this case, the apparently increased $H$, in Figure 6, in T13.1 $H$ values in the $\tau$ ranges below 200 s could represent a comparable upshift in the contribution of components whose relaxation, in the wild type, lies below the lower limit of the spectrum’s reliable range, rather than representing a downshift from $\tau$ approximately 1,000 s, as suggested in the preceding paragraph.

The spectral difference between line T7.2 and the wild type is smaller than that between T13.1 and the wild type. However, this is not really surprising, because although T7.2 is reduced in RG-I arabinans by approximately 70%, potato RG-I contains only a quarter as much arabinan as galactan, so this reduction amounts to a less than 2% change in wall composition (Skjøt et al., 2002).

Judging from the error bars in Figures 5 and 6, in T7.2, the rheological contribution of elements with $\tau$ values between 10 and 200 s is significantly greater than in the wild type, while Figure 6 suggests that in T7.2, the contribution of elements with $\tau$ values between 0.3 and 2 s is significantly reduced. This suggests that the removal of arabinan side chains from RG-I backbones in T7.2 may have increased, by some 10- to 100-fold, the viscous resistance that retards the movement of some wall component(s). If this component were RG-I itself, this would diverge from the above-mentioned expectation that side groups tend to restrict backbone mobility, but it would agree with the conclusion of Jones et al. (2003, 2005) that removing arabinans stiffens stomatal guard cell walls.

We suggested earlier (Ulvskov et al., 2005) that a difference in wall water status between the wild type and the transformant might contribute to the observed wall mechanical effects. Evered et al. (2007) demonstrated that hydration state influences primary wall mechanical properties. Tang et al. (1999) found that hydration greatly increased the mobility of pectic components of potato and water chestnut (Trapa spp.) cell walls. Moore et al. (2008) suggested that arabinans can be especially important to the mobility or “plasticizing” of pectins during water stress. Thus, the spectral shift toward a higher $\tau$ range in T7.2 might reflect a decrease in wall hydration, and thus in pectin mobility, when its arabinan content is reduced.

CONCLUSION

The BayesRelax algorithm for deducing relaxation spectra from rheological measurements has here been tested in what may be considered a very challenging or even worst-case scenario on biochemically active, turgid tissues, yet it was able to discriminate between wild-type and transgenic tissues with rather small wall changes. The computer program is of course equally useful for determining the relaxation spectra of materials that have been subjected to pretreatments that render them more amenable to frequency sweep measurements or even for solutions of isolated biopolymers that can be analyzed similarly.

MATERIALS AND METHODS

Plant Material

Wild-type potatoes (Solanum tuberosum 'Posmo'; Kartoffelforædlingstationen i Vandet) and the transformed lines T13.1 with reduced RG-I linear $\beta-1,4$-galactan (Sørensen et al., 2000) and T7.2 with reduced $\alpha$-1,5-arabinan (Skjøt et al., 2002) were grown in open air in containers in two-thirds peat moss and one-third Perlite. Tubers approximately 6 to 8 cm long were used for rheological measurements within 5 h of harvest. Samples were collected, and measurements made, over two growing seasons. Cylinders, 15 mm in diameter, were excised with a cork borer, and discs 3 mm thick were sliced from them using a custom-built potato guillotine (Mikrolaboratoriet). No more than two discs were cut from any one cylinder.

Rheological Measurements

Instrumentation

Small-amplitude oscillatory rheological measurements were performed using a Bohlin C-VOR rheometer (Malvern Instruments). The rheometer was equipped with a Peltier element to control the temperature (20°C ± 1°C) during the measurements. The measurement system consisted of two parallel, serrated plates (diameter, 15 mm). The discs of potato tissue, described above, were mounted between the plates before the measurement was started. During a 1-min equilibration period, the instrument was set to aim for a target gap of 2.80 mm between the plates and a NF of 30 g between the plates and the tissue disc. During the sequence of measurements that were then made on a particular disc, plate separation remained constant while NF declined. At the end of each measurement, the rheometer reported $G'$, $G''$, and NF values, this last being the value that prevailed at the moment the measurement ended.

Strain Sweep

A strain sweep was performed in order to determine the linear elastic range of the material. The small-amplitude oscillatory rheological properties were measured over the strain interval $10^{-3}$ to $10^{-1}$. 

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Frequency Sweep

For these measurements, maximum shear stress set at 26 Pa was used, since this gave strains within the linear range. Oscillation frequencies, 18 in all, over the range 0.001 to 10 s⁻¹, were applied successively to individual potato tissue discs in sets that kept the duration of the entire measurement run for any one disc to 20 min or less (15 min or less in runs that did not include the lowest frequency, 0.001 s⁻¹). At least 10 replicate measurements at each frequency, made on different individual discs, were ultimately recorded for each potato genetic line, except for the lowest frequencies, 0.001 and 0.002 s⁻¹, which require the longest measurement times and for which five and eight measurements, respectively, were recorded. Ten oscillations were averaged for the fastest frequency, five at 4 s⁻¹, two at 1.67 s⁻¹, and one oscillation for slower measurements, respectively, were recorded. For any one disc, frequencies were ordered in a randomized sequence to ensure that no single frequency was always recorded last, at a time when the material had undergone considerable NF relaxation, or first, when the NF was at its maximum. Data from tissue discs whose NF dropped nearly to 0 during measurement, indicating that the disc was no longer firmly clamped, were discarded.

Fitting Relaxation Spectra

Writing \( \omega \) for the frequency of oscillating strain, the relation between the relaxation spectrum \( H(\tau) \) and the measured storage and loss moduli \( G'(\omega) \) and \( G''(\omega) \), respectively, is given by the two Fredholm integral equations (Tschogel, 1989):

\[
G'(\omega) = G_0 + \frac{1}{\pi} \int H(\tau) \frac{\omega^2 \tau^2}{1 + \omega^2 \tau^2} \, d\tau
\]

\[
G''(\omega) = \frac{\omega \tau}{1 + \omega^2 \tau^2} \int H(\tau) \, d\tau
\]

For a viscoelastic liquid, the residual modulus \( G_0 = 0 \), and for a viscoelastic solid, \( G_0 \neq 0 \).

The quality of a fit to the data is measured through the \( X^2 \), which is defined in the conventional manner (i.e. for measurements at M frequencies):

\[
X^2(H) = \sum_{m=1}^{M} \left( \frac{G_m(\omega) - G_m(\omega_i)}{\sigma_i} \right)^2 + \sum_{m=1}^{M} \left( \frac{G_m(\omega) - G_m(\omega_i)}{\sigma_i} \right)^2
\]

where, leaving out primes, \( G_m(\omega) \) is the measured modulus, \( G_m(\omega_i) \) is the calculated modulus (from the integral equations), and \( \sigma_i \) is the SD of the noise at data point \( i \).

The estimation of \( H(\tau) \) using the \( X^2 \) may lead to a number of statistically acceptable but quite different solutions, as mentioned in the introduction. This problem is solved by regularization, which replaces \( X^2 \) by the functional (Tikhonov and Arsenin, 1977) where \( \alpha \) is a Lagrange multiplier and

\[
S(H) = \int [d^2H(\tau)/d(\ln \tau)^2] / H(\tau) d(\ln \tau)
\]

gives preference to smoother solutions. The functional form for \( S(H) \) may differ (Honerkamp and Weese, 1989), but the objective of the smoothness regularization functional \( S(H) \) used here is to impose an additional smoothness constraint upon the fitting procedure. As an alternative, \( S(H) \) may be chosen to give bias toward the baseline 0, but this choice can lead to additional (possibly artificial) peaks in the estimated spectrum (Hansen, 2008), as illustrated by the results obtained using the maximum entropy ("Masent") method of regularization (Fig. 7).

For a given choice of Lagrange multiplier, minimizing \( \alpha S(H) + X^2 \) will select the smoothest solution for the relaxation spectrum \( H(\tau) \) corresponding to the noise level in the data (scatter of individual measurements around their means), which determines \( \alpha \).

As \( S(H) \) takes its least value for a uniform \( H(\tau) \), this means that in the absence of constraints from the data, the estimate for \( H(\tau) \) will be a uniform (flat) function. The estimation of the value for the Lagrange multiplier has been given considerable attention in the literature (Honerkamp and Weese, 1990). A probabilistic (Bayesian) approach to the problem has been used in this article (details are given in Hansen, 2008).

Measuring the storage and loss moduli over the \( \omega \) interval 0.001 to 10 s⁻¹ allows the estimated relaxation spectrum to be interpreted safely over the corresponding range of \( \tau \) values 0.5 to 200 s (Davies and Andersen, 1997). Due to the correlation between neighboring points, induced by the smoothness constraint, this interval may be extended to \( \tau \) range of 0.1 to 1000 s, which corresponds to the conventional reliable interval used for interpretation. For numerical reasons, at least one extra decade of \( \tau \) values should be included, in the analytical procedure, at each end of this interval. The end points for the spectra were here chosen as \( \tau_{\min} = 0.001 \) s and \( \tau_{\max} = 100.000 \) s [below \( \tau_{\min} \) and above \( \tau_{\max} \) we assumed that \( H(\tau) = 0 \)]. The continuous distribution of \( \tau \) values in the Fredholm equations was approximated using 100 discrete \( \tau \) values distributed log linearly over that interval. These \( \tau \) values are plotted on the abscissa of Figures 5 and 6.

The 67% probability error bars were obtained by a calculation, specified in the BayesRelax program, that assumes (1) a Gaussian distribution of data error and (2) that the Fredholm equations correctly model the relaxation process. It deduces a probability distribution for each point in a spectrum using the sum of the \( \chi^2 \) and the \( \alpha^2 \) terms mentioned above (for an analysis of the probability problem that is involved, see Hansen and Wilkins, 1994).

BayesRelax can be accessed through its Web interface at www.BayesRelax.org. All spectra were calculated using the default settings, except that 100 points were calculated (specified under "optional parameters"). Normalization to NF of 30 g was carried out by two operations. The rheometer software had chosen to apply a series of \( \omega \) values, some members of which did not always correspond perfectly in different runs but, when not identical, did not differ greatly (closely adjacent \( \omega \) points in Fig. 3, A–C). Therefore, for normalization, data from \( \omega \) values that differed by less than 10% were pooled, and to this pool the mean of these \( \omega \) values was nominally assigned. Linear regression against NF was then performed, for every nominal \( \omega \) (whether from pooled, closely similar \( \omega \) values or not), on the pooled data from several potato lines (not just the three lines presented here) in order to avoid bias in the normalization. Each regression yielded a slope, \( \beta \), for the dependence of either \( G' \) or \( G'' \) on NF at each \( \omega \). Normalization consisted of adding the term \( (30 - NF) \beta \) to each \( G' \) or \( G'' \) measurement at a particular NF, using the \( \beta \) for the particular modulus and the measurement’s nominal \( \omega \).

Supplemental Data

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

Supplemental Data S1. An in-depth discussion of the technical issues listed in Table I.

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CORRECTIONS
Vol. 155: 246–258

Hansen S.L., Ray P.M, Karlsson A.O., Jørgensen B., Borkhardt B., Petersen B.L., and Ulvskov P. (2011) Mechanical Properties of Plant Cell Walls Probed by Relaxation Spectra.

The authors regret that problems with mathematical typesetting resulted in errors in the “Materials and Methods” section, beginning with the subhead Fitting Relaxation Spectra on p. 256 until the paragraph beginning “BayesRelax can be accessed through its Web interface at www.BayesRelax.org.” The corrected portion of the Materials and Methods is published below. In addition, the online supplemental data have been updated.

Fitting Relaxation Spectra

Writing ω for the frequency of oscillating strain, the relation between the relaxation spectrum H(τ) and the storage and loss moduli G′(ω) and G′′(ω), respectively, that are measured over a range of ω in a frequency sweep, is given by two integral equations called the Fredholm equations (Tschoegl, 1989):

\[ G'(ω) = G_0 + \int_0^\infty H(τ) \frac{ω^2τ^2}{1+ω^2τ^2} \frac{dτ}{τ} \]
\[ G''(ω) = \int_0^\infty H(τ) \frac{ωτ}{1+ω^2τ^2} \frac{dτ}{τ} \]

For a viscoelastic liquid, the residual modulus G_0 = 0, and for viscoelastic solids such as cell walls, G_0 ≠ 0.

The BayesRelax program searches for H(τ) functions that are compatible with a frequency sweep’s G′(ω) and G′′(ω) data by calculating from the Fredholm equations, for each trial H(τ) function, G′(ω) and G′′(ω) values over the range of ωs involved in the frequency sweep, and comparing these with the values that were actually measured. A best fit is indicated by a minimum value for χ^2, which is defined in the conventional manner, i.e. for measurements at M different frequencies:

\[ χ^2(H) = \sum_{i=1}^{M} \frac{(G'(ω_i) - G_m'(ω_i))^2}{σ_i^2} + \sum_{i=1}^{M} \frac{(G''(ω_i) - G_m''(ω_i))^2}{σ_i^2} \]

where, leaving out primes, G_m(ω_i) is the is the measured modulus, G(ω_i) is the modulus calculated from the appropriate Fredholm equation using the H(τ) function that is being tested, and σ_i is the SD of the measurements that were made at frequency ω_i. However, as mentioned in the introduction, a number of statistically acceptable but quite different H(τ) solutions may be obtained when minimum χ^2 is used as the sole criterion. This problem is solved by regularization (Tikhonov and Arsenin, 1977), which substitutes, for χ^2, the functional

\[ αS(H) + χ^2 \]

where α is a Lagrange multiplier, and

\[ S(H) = \int \frac{[d^2H(τ)/d(\ln τ)^2]^2}{H(τ)} d(\ln τ). \]

The objective of the S(H) just specified and, used here, is to impose a “smoothness” constraint upon the fitting procedure, in that minimizing this S(H) gives preference to smoother solutions. S(H) takes its least value for a uniform (flat) H(τ), so in the absence of constraints from the data, H(τ) would simply be flat.

Choosing, for the relaxation spectrum, the H(τ) function that minimizes αS(H) + χ^2, selects the smoothest solution that is compatible both with the mean values of the G′(ω) and G′′(ω) data and with the noise level (σ values) in these data, with which α increases, and with which χ^2 instead decreases as shown by the equation for χ^2 given above. The manner in which a value for α is specified has been given considerable attention in the literature (e.g. Honerkamp and Weese, 1990). A probabilistic (Bayesian) approach to this is used in the present paper, the details of which are given by Hansen (2008).

Forms of S(H) alternative to that given above could be used (Honerkamp and Weese, 1989). For example, an S(H) could be chosen that would give a bias towards the base line 0 (Elster and Honerkamp, 1991). However, this choice can lead to additional, possibly artifactual, peaks in the estimated spectrum (e.g. Hansen 2008), as illustrated by the results obtained using the maximum entropy (“Maxent”) method of regularization (Fig. 7).

Measuring the storage and loss moduli over the ω interval 0.001 to 10 s^{-1} allows the estimated relaxation spectrum to be interpreted safely over the corresponding range of τ values of 0.5 to 200 s (Davies and Andersson, 1997). Because of the correlation between neighboring points, induced by the smoothness constraint, this interval may be extended to a τ range of 0.1 to 1000 s, which corresponds to the conventional “reliable interval” used for interpretation. For numerical reasons, at least one extra decade of τ values should be included, in the analytical procedure, at
each end of this interval. The end points for the spectra were here chosen as $t_{\text{min}} = 0.001$ s and $t_{\text{max}} = 100,000$ s [below $t_{\text{min}}$ and above $t_{\text{max}}$ we assumed that $H(t) = 0$]. The continuous distribution of $t$ values in the Fredholm equations was approximated using 100 discrete $t$ values distributed log-linearly over that interval. These $t$ values are plotted on the abscissa of Figures 5 and 6.

The 67% probability error bars in Fig. 5 and 6 were obtained by a calculation, specified in the BayesRelax program, that assumes (1) a Gaussian distribution of data error and (2) that the Fredholm equations correctly model the relaxation process. It deduces a probability distribution for each point in a spectrum using the sum of the $\chi^2$ and the $\alpha_s$ terms mentioned above (for an analysis of the probability problem that is involved, see Hansen, 1994).
Problems in Frequency Sweep Measurements of Stress Relaxation in Cell Walls of Living, Turgid Plant Tissues

Peter M. Ray

This is an online addendum to Mechanical properties of plant cell walls probed by relaxation spectra by SL Hansen, PM Ray, AO Karlsson, B Jörgensen, B Borkhardt, BL Petersen and P Ulvskov. It discusses in some depth the physiological issues listed in Table 1 of the main paper.

We consider here phenomena that can cause turgor pressure (P) in potato tissue disc cells to change under the conditions of frequency sweep rheological measurements. These include (A) processes that could influence relaxation spectra directly, by being induced by the oscillating strain that the rheometer imposes to make modulus measurements, and (B) processes that could alter cell P over the course of the ca. 20 min measurement period that we employed. Because a plant tissue's elastic rigidity varies with P (Falk et al. 1958, Nilsson et al. 1958), processes of type (B) would affect the G' and G" values that the rheometer reports for any given oscillation frequency. This appears to be principally responsible for the large variation in G' and G values that occurred in our measurements. These variations correlated (main paper Fig. 4) with a progressive decline, during any measurement run (Fig. 1), in the compressive "normal force" (NF) between the tissue specimen and the rheometer's pressure plates, which clamp and hold it for oscillatory straining.

Issues and problems similar to those discussed here will probably affect rheometer measurements made on other kinds of tissues; the details that need consideration will depend on the form of the tissue fragment used, its anatomical structure, and the type of rheometer and tissue clamping method. The present analysis for potato tissue discs and the rheometer that we used will provide, we hope, an introduction to what needs to be considered for other tissues and instruments.

A. Processes that could contribute directly to relaxation spectra.

1. "In vivo stress relaxation". In the absence of an absorbable water supply (as with the tissue discs in our rheometer), the walls of turgid cells that are capable of growth can be expected to undergo a stress relaxation that is related to the irreversible extension that they undergo when the cells can take up water. Such a relaxation could also be provoked by the artificial strain that the rheometer imposes. This relaxation would be additive to that from reversible, viscoelastic relaxation processes in the walls, and could in principle yield its own relaxation-spectral peak whose position, on the relaxation time
(τ) scale, depends upon the cells' normal growth rate and their walls' elasticity (Cosgrove 1985, 1987).

Isolated discs of potato tuber tissue in water grow only slowly (~4% volume increase in 48 h) if incubated without auxin (Hackett & Thimann 1952). By the principles of in vivo stress relaxation (Cosgrove 1985), this growth rate would correspond to a relaxation time of about $6 \times 10^5$ s, far outside the range of our relaxation spectra. However, Hackett & Thimann's potato discs were pretreated in water for 24 h before growth measurements, to take them beyond the "initial phase" post cutting (Brauner et al., 1940), a phase that could involve more substantial irreversible cell enlargement. We therefore checked gravimetrically the short-term growth behavior of potato discs immediately after cutting (as were employed for our rheometer measurements). Weighed discs similar to those used in the rheometer were incubated in water or in 0.1 or 0.2M sucrose, followed, after either 20 or 120 min, by 0.28 M sucrose which was osmotically strong enough to just restore their initial $P$ and thus reverse the elastic expansion that they undergo in water (freshly cut potato discs neither gained nor lost water when placed immediately into 0.28 M sucrose, so equilibration with this medium, after exposure to a medium of higher $Ψ$, would restore their initial $P$). Re-weighing showed that they enlarged irreversibly at a rate of about 3% h$^{-1}$ in water and about 1.5% h$^{-1}$ in the 0.1M medium (these rates were recorded after both 20 and 120 min in these media), and hardly at all in 0.2M. (These are maximum possible rates of irreversible expansion, because part of the measured expansion could have been due to a possible hysteretic component of the preceding elastic expansion in the higher-$Ψ$ media.) This behavior corresponds to a $τ$ of about $1.6 \times 10^4$ s. This lies well outside the reliable range of our spectra and thus apparently could not be responsible for either of its principal peaks.

From a creep test (under a shear stress) performed on live potato tissue, Alvarez & Canet (1998) inferred a "Newtonian" irreversible strain component to the material's creep (i.e., a simple viscous flow). From the viscosity they inferred for this and the tissue elastic modulus that they reported, one can calculate that in a stress relaxation experiment it would yield a component with a $τ$ of about 100 s, similar to the higher-$τ$ peak in our potato stress relaxation spectra. But if such a component actually existed in potato cell walls it should have allowed, in the abovementioned tests that we made on potato discs, a rate of irreversible cell enlargement some 100 times what we actually measured. Furthermore, for multiple reasons which it is impracticable to enumerate here, the creep time course that Alvarez & Canet (1998, Fig. 7) actually published does not, in our opinion, support the existence in potato tissue cell walls of a capacity for irreversible
deformation, of anywhere near the magnitude that they inferred. Consequently we discount Alvarez & Canet's "Newtonian" retardation component as an explanation for the $\tau \sim 100$ s peak in our stress relaxation spectra.

2. Simple osmotic relaxation. Intuition might suggest that the cells' $P$ would increase as a result of the strain imposed by the rheometer. This would bring $P$ (and the cells' water potential, $\Psi$) above the value that exists at the osmotic equilibrium that prevails, in the absence of mechanical perturbation, between the cell contents and the cell wall. An osmotically driven (by the now-prevailing $\Delta \Psi$) relaxation of $P$ toward its equilibrium value would follow, and since the elastic modulus of a plant tissue varies with $P$ (Falk et al. 1958; Niklas 1988; Davies et al. 1998), the rheometer would register this as a mechanical relaxation. Its dynamics should be the same as those of the relaxation after a step change in $P$ imposed in the pressure-probe technique (Hüsken et al. 1978, Zimmerman & Hüsken 1979). For potato cells a half-time of 5 s has been reported (Michael et al. 1997, Table IVB); this corresponds to a $\tau$ of about 7 s. This falls right at the low point between the principal peaks in our relaxation spectrum (Fig. 5-6 of main paper), so apparently could not be responsible for either of these peaks.

Moreover, simple osmotic relaxation actually appears not to affect the present dynamic measurements at all, because the oscillating strain that the rheometer imposes is a shear strain, which does not change a material's volume nor, therefore, the cells' $P$. However, it could be a problem for the method by which potato discs are clamped between our rheometer's plates for measurements, to be discussed below, for which some basic considerations regarding strain-induced $P$ changes in cells need to be given.

**Strain-induced pressure changes in cells.** The $P$ in a pressurized, liquid-filled chamber changes if an externally imposed strain tends to change the chamber's volume, because the contained liquid volume cannot change (at least immediately) so its $P$ must change just enough to prevent the change in chamber volume ($\Delta V$) that would otherwise occur (Hepworth & Bruce 2000). This latter $\Delta V$ will tend to occur under an imposed uniaxial strain if the Poisson's ratio of the chamber's wall material (the ratio of its transverse contraction [or expansion] to lengthwise extension [or compression] under a uniaxial tensile [or compressive] stress) is less than 0.5 (as is true of most materials, including cell walls). An imposed compressive strain then tends to decrease the chamber's volume, resulting in an increase in $P$. An imposed extension strain tends to increase the chamber's volume, resulting in a decrease in $P$ if the chamber's contents are already under $P$, as are a plant cell's contents.
We verified, using a model spherical cell (a water-filled, pressurized, spherical rubber balloon coupled to a simple hydraulic manometer), that the $P$ within it increased when it was subject to uniaxial compression and decreased under uniaxial extension, but did not change at all when the model was deformed by shear. This is reasonable since shear strain, as noted above, does not cause a change in volume.

B. Changes in $P$ when tissue is clamped between rheometer plates

To hold a tissue disc between the rheometer's plates firmly enough that shear strain could be imposed on it without its surface slipping between the plates, these have to apply a compressive force to the specimen (the normal force or NF). This compression must increase the cells' $P$ (and $\Psi$) above its pre-existing, equilibrium level, so as explained above, an osmotic relaxation must then occur, involving efflux of water through each cell's plasma membrane into its cell wall, until the cell contents and walls regain water-potential equilibrium. This relaxation would show itself as a relaxation of the NF between the tissue disc and the pressure plate surfaces. Since, as noted above, the elastic moduli of a plant tissue depend strongly on $P$, relaxation of $P$ would decrease the measured $G$ values, which could cause them to correlate with NF, as in our published paper's Fig. 4.

1. Simple osmotic relaxation. The NF of about 100 g that develops initially (during automated advance of the movable rheometer plate in the first ca. 15 s of our 60 s pre-measurement "equilibration" period) corresponds to a stress of about 5.7 kPa on the disc as a whole. In the cell model experiments mentioned above, at small strains an externally imposed compressive stress increased $P$ by about 1/3 of the imposed stress' value (imposed force divided by the entire model's cross-sectional area). For potato cells the increase would probably be greater, because cell wall material has a smaller Poisson's ratio than rubber does, but in the disc tissue as a whole (not its surface cells, which will be considered shortly) the increase in $P$ could not exceed the over-all imposed stress, because the rheometer plates lack any mechanical advantage comparable to that of a narrow piston and cylinder in a hydraulic system.

By the classical osmotic method referred to parenthetically above in section A.1, combined with a freezing-point determination of tissue osmolarity, we determined the average $P$ of potato cells to be about 0.38 MPa, whilst Michael et al. (1997) reported a $P$ of 0.48 MPa from $P$-probe measurements on potato cells. Thus the initial NF could cause at most a 1-2% increase in the average cell's $P$. Since potato tissue's elastic modulus changes by about 3 times a change in its $P$ (Nilsson et al. 1958), the initially imposed NF could have caused at most about a 5% increase in its $G$ values. If the cells relaxed, osmotically, fully down to their previous osmotic equilibrium $P$, a 5% decrease in $G$
values could at most occur. But the actual variation in \( G \) values that correlates with NF greatly exceeded 5\%, so simple osmotic \( P \) relaxation resulting from compression of the tissue by the NF cannot account for most of the observed variation.

Furthermore, as noted above, this type of \( P \) relaxation should have a \( \tau \) of about 7 s, and thus should actually have gone virtually to completion within the 60 s equilibration period that preceded our first rheometer measurement with each disc. In agreement with this, Alvarez & Canet (1998, Fig. 6a) reported that compression of turgid potato tissue was followed by a (probably osmotic) stress relaxation with a half-time of about 4.2 s. So even had the compressive increase in \( P \) been larger than estimated above, its relaxation should not have influenced the measurements substantially.

Contrary to expectation for simple osmotic relaxation of \( P \), a rapid relaxation of NF often continued, beyond our 60 s equilibration period, for up to 100-200 s more. This was followed by a much slower relaxation "tail" that continued throughout the ca. 20 min duration of the tests on any one disc (Fig. 1).

2. **Ordinary viscoelastic relaxation.** The early, rapid NF relaxation must be due at least partly to whatever process is responsible for the longer-\( \tau \) peak at about 100 s in our relaxation spectra (main paper, Figs. 5 and 6). However, because this relaxation's magnitude could be at most only a fraction of the stress imposed on the cells by the NF, and this stress was, as noted above, small compared to the cells' \( P \), this relaxation could not cause the large effect on \( P \) that is needed to explain the large variations in \( G' \) and \( G'' \) with NF.

3. **Cell compression and collapse by pressure plate knurls.** The surfaces of our rheometer's pressure plates, between which a tissue disc is mounted in the instrument, are knurled to counteract a tendency of a tested specimen to slip, relative to the plates, as one of them moves relative to the other during a measurement. The knurls, or outwardly projecting pyramidal steel bumps, occur in an orthogonal array of rows 0.5 mm apart in both directions. They are about 0.15 mm high, above the bottoms of the grooves between the rows, into which grooves the sides of the knurls slope linearly. Since the average diameter of potato cells is about 130 \( \mu \)m (Michael *et al*. 1997, Davies *et al*. 1998), at a target plate separation of 2.8 mm between the tips of the knurls (see Methods) they should penetrate into each side of a 3 mm thick tissue disc by almost one cell diameter, severely compressing or collapsing the impacted cells. Rows of depressions visible on both surfaces of the discs after a set of measurements was completed, showed that cell compression or collapse did occur. Besides the cells that are directly impacted by the tips of the knurls, adjacent surface cells out to a distance of about 0.13 mm from the knurls'
tips must become strongly compressed. From the numbers just given one can estimate that about 25% of the surface cells on either side of a potato disc must become compressed by 50% of their initial volume or more. Compression would also strongly affect at least the second layer of cells beneath the directly impacted ones, since potato tissue contains a negligible fraction of intercellular air channels that could absorb the compression that progresses inward from the disc's surface as the pressure plates advance.

The initial compression of these cells by the knurls must generate a large stress on them, with a large increase in their $P$, probably exceeding their own initial $P$. This would probably not rupture their cell walls, since the walls can support at least twice the cells' normal (within the tuber) $P$ when potato tissue is placed in water and osmotically equilibrated with it. The severely compressed cells must instead rapidly lose, osmotically, most of their water and shrink greatly, probably over a period of a few hundred s at most. This collapse is probably largely responsible for the initial, rapid phase of NF relaxation noted earlier. Water made available to other cells, in their common cell wall space, by collapse of the directly impacted ones, would be expected to raise $P$ in parts of tissue remote from the knurl impact areas, since their initial $P$ of about 0.4 MPa lies well below that required for osmotic equilibrium with water (about 0.7 MPa, the cells' average solute osmotic potential). This rise would increase the tissue's rigidity and contribute to the unexpectedly high initial values of NF (far above its 30 g pre-set target value) that were recorded, after the first measurement that was made, in many of the measurement runs (Fig. 1).

4. Solute leakage from cells impacted by pressure plate knurls. Water loss from cells that are compressed or collapsed by the knurls must raise these cells' internal solute concentration far above normal. This will very likely lead to substantial solute leakage through their plasma membranes into the cell wall space, both as a result of membrane damage that they will probably suffer during their sudden compression, as well as simply high-concentration-driven efflux. As these solutes diffuse into nearby cells' wall space, lowering its water potential, this will force these latter cells to lose water osmotically and reduce their $P$. This effect would gradually extend, by solute diffusion, throughout the entire disc after surface and subsurface cell compression or collapse is completed. The $P$ of 1 x 1 mm thick potato tissue strips equilibrated osmotically, after a change in external mannitol concentration, with a half-time of about 3 min (Virgin 1955), in a process that was doubtless governed by solute diffusion through the tissue's cell walls (free space), since it was much slower than the osmotic equilibration of potato cells after a step change in their $P$ in a pressure probe apparatus as noted earlier. For our potato discs this half-
time should be more than doubled, because our tissue was 3 times as thick as Virgin's, and solute would have diffused into our discs' interior from just 2 sides rather than from their strips' 4 sides; cell solutes' weighted mean diffusion coefficient may well also be smaller than mannitol's. The resulting decline in $P$ would thus be expected to continue throughout the 20 min measurement period. It would contribute to the gradual decline in NF and $G$ values that occurred during most of the 20 min measurement period, after the rapid initial NF relaxation, as noted above.

A rough estimate of the probable magnitude of this effect is as follows. From their diameters and approximately isodiametric shape one can estimate that the cells that must be severely compressed on both sides of a disc by the pressure plate knurls constitute at least 2% of the disc's volume. If most of the solutes in these cells were to enter, and diffuse throughout, a free space amounting to perhaps 5% of the remaining tissue volume, the osmolarity in that space would increase (above whatever its initial value was) by about 40% of the initial internal osmolarity of the cells. This should cause the $P$ in cells throughout the rest of the disc to fall greatly. In the actual situation the directly collapsed cells might not leak all their solutes, but indirectly compressed or distorted cells beneath and to either side of them would probably leak some. It seems clear that a substantial decrease in $P$ and thus in measured $G$ values should occur, since at their normal $P$ about half of potato cells' rigidity has been estimated to be due to their $P$ (Davies et al. 1998).

In addition, it seemed possible that solute leakage into the free space from cells throughout the disc, provoked by the compressive stress that the pressure plates impose on it, or by the (very small) shear strains to which it is subjected during the measurements, might occur and cause $P$ to fall. However, tests we made of discs placed under even a continuous 100 g compressive load, applied through a flat surface, indicated that their $P$ did not fall any faster than that of unloaded discs, and indeed no faster than could be explained by evaporative water loss from their edges. Therefore, this possibility seems contra-indicated.

In about a quarter of our rheometer runs, the NF that was recorded at the end of the first $G'$ and $G''$ measurement, at less than 100 s after the equilibration period, was less than 30 g, and only the second, slow phase of NF relaxation was seen over the length of the run, so the initial, rapid NF relaxation phase was evidently missing. These may be cases in which the disc thickness (see item C, below) was sufficiently less than 3.0 mm that, in advancing toward the target gap of 2.8 mm, the pressure plates' knurls did not penetrate into the disc far enough to cause substantial collapse of its surface cells. If so,
the following two mechanisms may have been responsible for the modest NF relaxation that was observed in these cases.

5. **In vivo stress relaxation.** The $P$ in the surface cells impacted by the pressure plate ridges very likely exceeded, initially, the potato cells' yield threshold, allowing some *in vivo* stress relaxation to contribute to the rapid NF decline early in a run. Since, according to the preceding analysis, the general $P$ in the tissue disc would have fallen below its initial (pre-clamping) value, and hence below the yield value\(^1\), by the time that the more gradual, post-100 s decline in NF was occurring, *in vivo* stress relaxation should not have contributed to the latter decline. Without knowing how far the initial (post clamping) $P$ rose above the yield threshold or how long it remained there, it is not possible to estimate how much of an effect this process could have had on the system.

6. **Evaporative water loss.** When clamped in the rheometer, both faces of a potato disc were protected, by the rheometer's pressure plates, from evaporation, but its edges were directly exposed to the air. From measurements that we made of evaporation from the edges of potato discs 3 mm thick into still air we calculate that a disc would have lost about 2\% in volume from its edges during the measurement period, corresponding to a decrease in $P$ of about 0.1 MPa. This is only a rough expectation, because atmospheric humidity, temperature and motion, which affect evaporation rate, would not have been the same in our evaporation tests as within the rheometer. But evaporative decrease in $P$ would evidently have contributed to the correlation of $G$ values with NF that we observed.

C. **Variations in initially imposed NF.** After a tissue disc was inserted into the rheometer, its movable pressure plate advanced toward the target NF (30 g) and plate separation (2.80 mm) settings over about 10 s in an intermittent, "searching" fashion. The first NF values reported in different measurement runs varied considerably (Fig. 1), some of them greatly exceeding 30 g. This indicates that the compromise between these targets that was reached, at the time that plate advance stopped, was not consistently repeatable.

Besides intrinsic, stochastic variations in the instrument's operation, a factor that likely contributed to initial differences in NF among discs was variation in initial disc thickness (some such variation was known to occur). This variation would obviously affect the NF value that would be reached if pressure plate advance were to stop at exactly the set target separation distance.

Initial variations in NF partially converged during the rapid decline period, early in measurement runs, in the sense that unusually high initial NF values declined more rapidly than initially low ones. However, appreciable differences in NF between samples
still remained at the onset of the slowly declining, "tail" phase that continued through the entire measurement period, and these differences persisted to the end of the measurement runs (Fig. 1). Thus, although changes in NF in the course of this decline probably correlate well with changes in P, an absolute relation between NF and P values in different runs cannot be expected. This is because the relation between the two depends on the uncontrolled actual area of contact and extent of penetration into the tissue, of the pressure plates' knurls (cf. Wei et al. 2001). Therefore variation, between runs, in the general level of NF values arising from initial differences in NF cannot be expected to correlate well with P or with reported G values. It would instead cause a scatter or imperfection in the correlation of G values with NF (main paper's Fig. 4) that would be additive to any biological variation in mechanical properties between different discs, plus any imprecision in the rheometer's measurement of G values, in contributing to the "noise" that would remain in the data after they have been normalized to a standard NF value.

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Fig. 1. Time course of changes in normal force (NF) after successive oscillating-strain measurements on different individual potato discs. Each curve pertains to a different disc. The NF values reported by the rheometer were for the end of each measurement, and are plotted in this figure for that time, subsequent to the start of the first measurement on the given disc (thus not including the standard 1 min equilibration period prior to beginning the first oscillating-strain measurement, see Methods in main paper). Successive measurements on any one disc involved different oscillation frequencies, in a random order. The measurements were on discs from wild-type tubers.