High-pressure cell for simultaneous dielectric and neutron spectroscopy

Alejandro Sanz,1,a),b) Henriette Wase Hansen,1,a),c) Bo Jakobsen,1 Ib H. Pedersen,1 Simone Capaccioli,2 Karolina Adrjanowicz,3 Marian Paluch,3 Julien Gonthier,4 Bernhard Frick,4 Eddy Lelievre-Berna,4 Judith Peters,4,5 and Kristine Niss1,a),d)

1Glass and Time, IMFUFA, Department of Science and Environment, Roskilde University, P.O. Box 260, DK-4000 Roskilde, Denmark
2Department of Physics, University of Pisa, 56127 Pisa, Italy
3Institute of Physics, University of Silesia, ul. Uniwersytecka 4, 40-007 Katowice, Poland
4Institut Laue-Langevin, CS 21356, 38042 Grenoble Cedex, France
5Université Grenoble Alpes, CNRS, LIPhy, 38000 Grenoble, France

(Received 29 September 2017; accepted 2 February 2018; published online 22 February 2018)

In this article, we report on the design, manufacture, and testing of a high-pressure cell for simultaneous dielectric and neutron spectroscopy. This cell is a unique tool for studying dynamics on different time scales, from kilo- to picoseconds, covering universal features such as the α relaxation and fast vibrations at the same time. The cell, constructed in cylindrical geometry, is made of a high-strength aluminum alloy and operates up to 500 MPa in a temperature range between roughly 2 and 320 K.

In order to measure the scattered neutron intensity and the sample capacitance simultaneously, a cylindrical capacitor is positioned within the bore of the high-pressure container. The capacitor consists of two concentric electrodes separated by insulating spacers. The performance of this setup has been successfully verified by collecting simultaneous dielectric and neutron spectroscopy data on dipropylene glycol, using both backscattering and time-of-flight instruments. We have carried out the experiments at different combinations of temperature and pressure in both the supercooled liquid and glassy state. Published by AIP Publishing. https://doi.org/10.1063/1.5007021

I. INTRODUCTION

Supercooled liquids are metastable with respect to the crystalline state and induced, e.g., by rapid cooling below their melting point, $T_m$, to avoid nucleation and crystal growth. Alternatively, compression can be utilized to create metastable liquids above a certain melting pressure, $P_m$. If temperature keeps decreasing (or pressure increases), the structural relaxation is found to be more and more hindered and eventually the rearrangement of the molecules becomes so slow that the equilibrium volume and enthalpy cannot be reached on experimental time scales. This marks the onset of the liquid-to-glass transition. Pressure is an important variable for disentangling temperature and volumetric effects on the slowing down of the dynamics when approaching the glass transition. In the proximity of the glass transition temperature, $T_g$, the typical time scale for molecular rearrangements is minutes or even hours. Along with this extremely slow dynamics, glass-forming materials present other universal features such as fast vibrational processes as well as fast relaxations. The slow structural relaxation freezes when liquids enter the glassy state while vibrations are still present. These fast vibrations can be interpreted as short time dynamics around the minima of the energy landscape. They are governed by the shape of the minima and present characteristic time scales of $\sim 10^{-12}$ s.

In amorphous solids, one also observes an excess in the vibrational density of states as compared to the Debye prediction for crystals. The Boson peak is considered to be a characteristic feature of disordered solids. It is now generally agreed that an appropriate comprehension of the glass transition must encompass fast and slow dynamics. In order to study the broad range of dynamics covering molecular movements from picosecond to kilosecond time scales, it is necessary to combine complementary techniques, such as dielectric and neutron spectroscopy.

For a better understanding of how slow and fast dynamics are correlated with each other at varying thermodynamic conditions, we present here a novel experimental setup to perform simultaneous dielectric and neutron spectroscopy. The cell, developed at Roskilde University in collaboration with the Institut Laue-Langevin (ILL), has been designed for a temperature range of approximately 2–320 K and for a maximum pressure of 500 MPa which is 5000 times atmospheric pressure. This setup is a unique tool for studying a variety of systems with dynamics on a large range of time scales, such as viscous liquids, polymers, and proteins. With the current design, the high-pressure (HP) cell is only suitable for samples with sufficiently low viscosity at room temperature. So, apart from molecular liquids, only low molecular weight polymers, oligomers, or macromolecular solutions can be loaded. By performing simultaneous dielectric and neutron spectroscopy experiments, we are able to monitor slow and fast (relaxational and vibrational) dynamics under exactly the same conditions, precluding unwanted external effects associated with different environments, for example, shape and material of the cell.

---

A. Sanz and H. W. Hansen contributed equally to this work.

Electronic mail: asanz@ruc.dk

Electronic mail: hansenhw@ill.fr

Electronic mail: kniss@ruc.dk
container and heat and pressure transfer. Achieving thermodynamic equilibrium is not trivial since proper transmission of pressure into viscous liquids and the equilibration of temperature in heavy sample holders are real problems. These critical issues may slightly affect the thermodynamic state of the sample and therefore modify its dynamic behavior, yet in the same way for our simultaneous experiments. This is relevant when one looks at the $\alpha$ relaxation near the glass transition, where small pressure and/or temperature variations strongly modify the characteristic relaxation time. Additionally, biological samples start to undergo unfolding between 200 and 400 MPa and it is therefore difficult to reproduce exactly the same conditions twice. The simultaneous measurements are especially important when the sample is not stable, for instance, due to physical or chemical processes such as aging, crystallization, chemical degradation, polymerization, or water uptake.

In the above-mentioned case studies, the possibility to apply pressure and temperature changes is of crucial importance since the volume reduction can induce a peculiar effect on the phenomena, and the time evolution, being irreversible, is very difficult to study on both fast and slow dynamics under the same conditions.

Studying simultaneously the slow (by dielectric) and fast dynamics (by neutron spectroscopy) on the same system undergoing physical aging after a jump is of paramount importance since the same experiment repeated on two separate setups is much less reliable. The relevance of physical aging in out-of-equilibrium liquids has been recently highlighted by Niss, who proposes the existence of isostructural lines in the equilibrium phase diagram and the possibility of understanding isobaric aging in simple liquids in the framework of isomorph theory.

A neutron diffraction cell for simultaneous dielectric and neutron measurements for crystallization studies was developed by Jiménez-Ruiz et al. to ensure that the measurements were done at exactly the same conditions, a critical issue in kinetics studies such as crystallization.

In dielectric spectroscopy (DS), the sample is placed between two conducting electrodes, creating a capacitor. This technique measures the complex capacitance $C^*$ as a function of the applied frequency $\nu$, typically from mHz to MHz. From the capacitance, the complex dielectric permittivity of the sample $\varepsilon^*(\nu) = \varepsilon'(\nu) + i\varepsilon''(\nu)$ can be calculated either through the capacitance of the empty capacitor or from knowing the geometry of the capacitor. When the material undergoes a relaxation process in response to an alternating electric field, the imaginary part of the dielectric permittivity, $\varepsilon''(\nu)$, referred to as the dielectric loss, shows a maximum. The position of this peak $\nu_{\text{max}}$ provides information about the characteristic relaxation time $\tau$ of the underlying dynamics through the relation $\tau = (2\pi \nu_{\text{max}})^{-1}$.

In incoherent neutron spectroscopy (NS) for samples with a large content of hydrogen, such as organic liquids, the incoherent dynamic structure factor, $S(Q, \omega)$, is measured as a function of the energy transfer, $\hbar\omega$, and momentum transfer, $Q$, and provides information on the self-motion of atoms. The inelastic signal in incoherent scattering for hydrogenous samples is often seen as a broadening of the elastic peak commonly referred to as quasielastic neutron scattering (QENS). The energy resolution of the neutron spectrometer determines the time scale of the experiment. Spatial information of the dynamics can be obtained through the dependence of the dynamic structure factor on the momentum transfer $Q = (k' - k)$, where $k$ is the wavevector of the incident neutron and $k'$ is the scattered wavevector. The momentum transfer $Q$ can be related to the scattering angle via $Q = (4\pi/\lambda) \sin(\theta/2)$, where $\lambda$ is the neutron wavelength and $\theta$ is the scattering angle. Neutron backscattering spectroscopy (BS) normally covers molecular motions between $10^{-9}$ and $10^{-10}$ s, while instruments based on time-of-flight (TOF) methods provide information at the sub-nanosecond regime. Features such as vibrations, fast relaxations, and the Boson peak have typically been detected by TOF spectrometers.

An advantage of combining dielectric spectroscopy and incoherent neutron scattering is that the optimal cell geometry is identical for both methods. The area should be as large as possible and the thickness of the sample should be small; for a cylindrical capacitor, which is used in this design, the dielectric signal scales with the height of the capacitor and is inversely proportional to the distance between the electrodes. For incoherent QENS, the sample thickness must remain small in order to minimize multiple scattering effects, while a large sample area enhances the signal.

With this new setup, the sub-nanosecond dynamics can be measured simultaneously with the slow $\alpha$ relaxation seen in dielectric spectra, a universal fingerprint of the glass transition. Given that the $\alpha$ relaxation stretches over several decades, it is still visible with DS when it enters into the NS window, at least as a high frequency tail.

In this paper, we report on the design, manufacture, and testing of a HP cell for carrying out dielectric and neutron spectroscopy simultaneously. This setup can be utilized in different neutron spectrometers in order to combine DS and NS at different energy transfers and energy resolutions, thus accessing different time scales, in broad temperature and pressure ranges. We present the simultaneous collection of DS and NS data performed on the spectrometers IN16B and IN6 (ILL) for different combinations of temperature and pressure, demonstrating the utility of the technique.

II. DESCRIPTION OF THE HIGH-PRESSURE CELL FOR SIMULTANEOUS DIELECTRIC AND NEUTRON SPECTROSCOPY MEASUREMENTS

An important requirement for constructing the cell is to reduce as much as possible the neutron attenuation. Materials with high tensile properties, low neutron absorption and incoherent scattering cross section are required. We have carried out finite element calculations for different prototypes in order to optimize the dimension, shape and type of material, and have measured the transmission and background on the spectrometers IN6, IN16B, and IN13 (ILL). Based on these results, the cell was constructed in cylindrical
geometry using the aluminum alloy EN AW 7049A-T6, which contains the following composition (mass %, balance Al): Zn 7.2-8.4, Mg 2.1-3.1, Cu 1.2-1.9, Fe 0.50, Si 0.40, Mn 0.50, and Cr 0.50. At 300 K, EN AW 7049A-T6 shows a yield stress of 530 MPa. Given its high content of aluminum (≈mass 90%), EN AW 7049A-T6 shows a low incoherent neutron scattering cross section and thus, a low background in the data.

The development of high-pressure equipment in neutron facilities has been the subject of intense research activity over the last decades (see, e.g., Refs. 19 and 24–27). The design of our device was inspired by the high-pressure cell recently developed by Peters and co-workers, in collaboration with the Service for Advanced Neutron Environment (SANE) at the ILL. Proposed designs utilized a piston or floating barrier to separate the sample volume from the pressure-transmitting liquid. Peters et al. also discussed the special requirements that a sample holder must fulfill for an optimal performance. For example, the sample stick, which was also designed at the ILL, is not only responsible for holding the sample but also for transmitting pressure onto the sample through a heated capillary that links the sample cell to a...
compressor outside the cryostat. The capillary must be thermally isolated as it passes the cold-point in the cryostat in order to avoid unwanted freezing of the pressure-liquid.29

Our HP neutron cell is designed according to similar principles; however, it integrates an additional component into the neutron beam; a cylindrical capacitor for measuring the dielectric properties of the sample. Within the bore of the HP container, there is a coaxial cylinder surrounding a rod. Both the hollow cylinder and the rod, the outer and inner electrodes of the capacitor, respectively, are also made of the aluminum alloy EN AW 7049A-T6. This is challenging compared to ordinary HP cells for neutron scattering since the electrodes have to be connected to the electronics through coaxial cables (RG178 with an outer diameter of 1.8 mm) which will be fed into the pressurized zone by a sealed plug with feedthroughs.

A. Main components of the high-pressure cell

The high-pressure DS-NS cell is presented in an exploded-view drawing (Fig. 1). A hollow cylindrical monobloc with an inner diameter of 12 mm is the main-body piece. The outer diameter is reduced as much as possible in front of the beam section to minimize the neutron attenuation and diffuse scattering (background). It is 32 mm at the center, leading to 10 mm wall thickness in the neutron beam, and 43 mm at the thickest parts. The top end is connected to a capillary where the pressure-transmitting liquid is injected. In order to preclude unwanted mixing of the sample with the pressure-transmitting liquid, we use a floating barrier that consists of a disc made of brass surrounded at its middle part by a rubber O-ring seal. A reservoir of sample volume separates the floating barrier from the dielectric capacitor. At the other end, copper wires are fed into the pressurized zone through a plug made of brass sealed with an epoxy encapsulant and connected to the capacitor. Optimum tightness of the plug is achieved by the action of two seals, one made of copper and the other one made of rubber for high- and low-pressure ranges, respectively. O-rings remain tight at low temperature/high pressure when they are pre-compressed in a groove of appropriate dimensions. At low pressure, the rubber O-ring will deform, thus keeping the sealing tight. In particular, we use the elastomer NBR 90 (M-Seals) which has an excellent resistance to mineral oils, aliphatic hydrocarbons, and many organic solvents and chemicals. The plug is held by a 22 mm long locking nut made of a copper-beryllium alloy. A cross-sectional view displayed in the left-top corner of Fig. 1 shows the location of cadmium foils for shielding the wires, sealed plug, floating barrier, and the thickest parts of the HP container.

Figure 2 shows a schematic representation of the sample geometry illustrating the location of electrode-electrode and cell-electrode sample volumes. We expect identical behavior for the sample occupying these two compartments given the same thickness and that the high-pressure container and the electrodes are both made of the same aluminum alloy. For neutrons, this results in a hollow cylinder sample geometry with a thickness of 0.3 mm.

B. Strengthening of the high-pressure cell

It is worth noting that compression may induce elastic and plastic deformation of the cell. The cell must operate in the elastic regime for optimal performance. In order to increase the maximum pressure at the elastic regime, the HP cell was autofrettaged by initial compression up to 530 MPa. The autofrettage process is performed by applying pressure above the elastic region. For a monobloc cylinder, according to the Lamé equations and the distortion energy theory for plasticity criterion,19 the bore of the cylinder will start to deform plastically at a pressure $P$.

$$P = \sigma_Y \frac{(K^2 - 1)}{\sqrt{3}K^2},$$  \hspace{1cm} (1)

where $\sigma_Y$ is the yield stress of the material and $K = r_o/r_i$ is the wall ratio ($r_o$ and $r_i$ are the outer and inner radius of the cylinder, respectively). By means of this strengthening process, it is possible to increase the maximum pressure at which the cell yields. In this procedure, the monobloc vessel is first bored with a slightly narrower inner diameter. Then it is pressurized above the limit defined by Eq. (1) in such a way that the material at the bore deforms plastically. Once the pressure is released, the material that was just deformed elastically will radially compress the area close to the bore that initially suffered plastic deformation. This raises the pressure limit at which the cell starts to yield. For a monobloc with $K > 2.2$, the pressure limit is given by

$$P_l = 2P.$$  \hspace{1cm} (2)

Equation (2) also determines the pressure for the autofrettage process. Finally, the bore is machined to achieve the desired diameter and the cell will operate in an elastic regime as long as the pressure limit $P_l$ is not exceeded.

C. Dielectric capacitor and plug with electrical feedthroughs

A detailed drawing of the cylindrical capacitor and sealed plug is presented in Fig. 3. The design of the capacitor constitutes a technical challenge since an optimum performance
at elevated pressure requires a minimum deformation of the device, avoiding mainly electrode collapse, given the fact that a physical contact between the electrodes would ruin the measurements. The outer electrode has an inner diameter of 11.4 mm (external diameter 11.7 mm), while the inner electrode has a diameter of 11.1 mm, leading to an internal gap for the sample of 0.15 mm. The inner electrode sits on the bottom insulating spacer which through a cup-like shape keeps the outer electrode at a fixed position ensuring the right sample gap. The top and bottom spacers are made of polyether ether ketone (PEEK). Holes in the spacers allow the connection of the electrodes to copper wires and these spacers also serve for insulating the capacitor from the HP container. The height of the capacitor is 24 mm. Simple calculation by applying Gauss’ law gives an empty capacitance of 50 pF, in close agreement with the value we found experimentally, approximately $55 \pm 1$ pF. This value may change slightly depending on the precise geometry of the capacitor and the exact temperature but can easily be determined with high precision before each experiment.

A key technical issue is how to feed the wires from outside into the pressurized zone and to connect them to the capacitor while preventing any leak at high pressure and keeping the capacitor electrically isolated from the rest of the cell. For this kind of mechanical sealing and electrical shielding, we developed a plug made of brass with a single cylindrical channel that guides the wires from outside into the cell. The remaining gap is filled with a dense paste that is then cured to transform...
The stick was designed and developed by SANE at the ILL. Its geometry was optimized for these high-pressure and dielectric measurements. The cell was connected to a neutron sample stick especially optimized for working at these low temperatures, pressure must be applied at higher temperatures and maintained upon cooling through the heated capillary. The high-pressure DS-NS cell was connected to a neutron sample stick especially optimized for these high-pressure and dielectric measurements. The stick was designed and developed by SANE at the ILL.

The sample stick plays three main roles; first, it allows locating the sample in the path of the neutron beam within the calorimeter of a cryostat; second, it drives the pressure-transmitting medium from the compressor to the sample through a capillary, avoiding freezing of the pressure-liquid at the cold point of the cryostat; and third, it brings two coaxial cables that are connected through the plug to the capacitor. We utilized as pressure controller the electrical pressure multiplicator Louise which transmits hydrostatic pressure to the sample with Fluorinert™, a non-hydrogenous liquid.

Two sensors measure the pressure in such a way that the controller may decide to increase or decrease the pressure in order to reach and maintain the desired set-point. The pressure controller Louise has also been developed at the ILL and can be operated with the instrument control software NOMAD.

The dielectric electronics used for measuring the capacitance of the sample consists of a commercial LCR meter (Agilent E4980A) covering the frequency range $10^2$–$10^6$ Hz, while a multimeter in combination with a frequency generator covers the low frequency window $10^{-3}$–$10^5$ Hz. This particular setup and the MATLAB-based control were developed at Roskilde University and uses an automatic switch to couple the setup at 100 Hz. In order to synchronize neutron and dielectric measurements, an interface of the dielectric software was integrated in NOMAD at the ILL.

A. Experimental details

In the following lines, we explain how the cell and capacitor are filled as well as the order of assembly. First, the separator or floating barrier is inserted into the high-pressure cell at room temperature and the bore of the HP vessel is then filled with enough liquid sample to ensure a complete fill of the cell after the introduction of the capacitor. For the dielectric insert, the capacitor is filled using capillary force; the gap separating the electrodes of the capacitor is simply filled by placing the assembled capacitor in the liquid sample and letting it get sucked inside by capillary effects. The insulating spacer of the capacitor possesses four gaps for making the filling of the capacitor possible. If the liquid is too viscous, this filling can be accelerated by moderate heating. The capacitor creates the electrode-electrode sample compartment. Once the capacitor is filled, it is inserted into the HP cell. The filled gap between the inner wall of the HP vessel and the outer electrode forms the cell-electrode sample compartment. Then, as illustrated in Fig. 1, the plug with feedthroughs incorporating rubber and copper seals is connected to the capacitor. The cell is finally locked with the nut and potentially trapped air is let out as it is closed. A schematic representation of the sample geometry is shown in Fig. 2.

B. Examples of simultaneous data collection in dielectric and neutron spectroscopy techniques

The investigation of glass-forming liquids is one of the major envisaged applications and the following example aims to show the advantage of simultaneous measurements without proceeding to a detailed quantitative evaluation of the science. We present simultaneous pressure and temperature measurements on the molecular glass former dipropylene glycol (DPG). This hydrogen-bonded liquid has a $T_g$ of 195 K at atmospheric pressure and a pressure coefficient $d^2T_gdp = 0.08$ K MPa$^{-1}$.

As an example, we report first on simultaneous neutron and dielectric pressure scans carried out on this sample on IN16B at the ILL and then on simultaneous dielectric-TOF experiments from IN6 (ILL). For DPG, if the pressure is changed between 0.1 and 400 MPa at 270 K, the density varies between 1.112 and 1.237 g/cm$^3$, which for the described sample geometry and the cross sections for DPG results for the neutron experiments in an average transmission of 0.66 (largely dominated by the incoherent scattering of hydrogen). We should stress that such a high scattering probability induces multiple scattering, which will influence a quantitative evaluation of the data, but it does not invalidate the qualitative behavior, as described below. The top panel of Fig. 4 displays a double logarithmic representation of the imaginary part of the capacitance of DPG as a function of frequency. On the low frequency side, we observe a power law contribution due to free charge conductivity which shifts towards lower frequencies as pressure goes up. On the high-frequency side, the low-frequency tail of the $\alpha$ relaxation also moves towards lower frequencies with pressure and finally the $\alpha$-peak enters the experimental frequency window. In the bottom panel of Fig. 4, we have selected two frequencies to illustrate how we can monitor the change with pressure in conductivity at 1 kHz and in the relaxation of the liquid at 100 kHz (right axis).
The reduced mobility with pressure produces a quasielastic signal narrows with the slowing down of about 60%. The inelastic intensity decreases accordingly as and were summed over Q\textsuperscript{ing instrument IN16B (ILL), normalized by the high pressure data point, of pressure at 270 K. Neutron intensities were acquired on the backscatter- and inelastic fixed window scans with neutrons (bottom), on DPG as a function FIG. 4. Simultaneous dielectric spectroscopy measurements (top), and elastic and inelastic fixed window scans with neutrons (bottom), on DPG as a function of pressure at 270 K. Neutron intensities were acquired on the backscattering instrument IN16B (ILL), normalized by the high pressure data point, and were summed over Q. The bottom panel also shows the pressure dependence of the sample capacitance (imaginary part) at two selected frequencies: 1 and 100 kHz (vertical dotted lines in the top panel). Empty symbols (top) and crosses (bottom) correspond to a second pressure scan carried out 15 h later.

to as IFWS (ΔE ≠ 0) and can be alternated with EFWS during temperature or pressure changes. Here are shown IFWS data for an energy offset of 2 μeV, together with the EFWS (left axis). The reduced mobility with pressure produces a progressive increase of the Q-integrated elastic intensity by about 60%. The inelastic intensity decreases accordingly as the quasielastic signal narrows with the slowing down of the α relaxation in the measured T, P-range for DPG and thus its spectral weight decreases at 2 μeV energy transfer by about 50%. While the size and slope of the observed relative changes might be influenced by multiple scattering, the observed qualitative trend clearly stems from first-order scattering.

Up and down pressure cycles were successfully carried out exhibiting reproducible values of the scattered neutron intensity and sample capacitance, corroborating that the floating barrier applies pressure to the sample and is able to retract again to the initial position when pressure is released. As an example, we compare the pressure dependence of dielectric and neutron data between two scans performed 15 h apart, illustrated with x in Fig. 4. One observes that the results are quite similar in both pressure scans.

C. Practical advantages

Studying materials under pressure in general requires large sample holders in comparison to standard ambient pressure setups, which makes temperature changes and waiting for the sample to reach equilibrium a relatively slow process. These waiting times for the sample to reach equilibrium can be monitored carefully in the dielectric signal. The α relaxation is one of the dynamic features that can be probed with dielectric spectroscopy. Due to its extreme sensitivity to pressure and/or temperature changes, the dielectrics acts like an internal probe to define exactly the thermodynamic state of the system. We take advantage of this powerful property to gain information on, for example, whether or not the sample is equilibrated when approaching the glassy state either by cooling or compression. We can also gain insight into potential difficulties in transmitting the pressure onto the sample. Figure 5 (top) shows the dielectric α relaxation of DPG when entering the glassy state at fixed P = 200 MPa upon cooling down to 210 K. The dielectric spectra from 10\textsuperscript{-1} Hz are roughly 10 min apart. By reaching the 210 K and 200 MPa state point on cooling at constant pressure (top panel, Fig. 5), one observes that the system is mostly thermally equilibrated with a continuous shift of the α relaxation towards lower frequencies. Finally, after approximately 4 h at IN6 (this time depends on the cooling rate of the cryostat and the heat conductivity of the sample environment), the peak of the α relaxation is located at the characteristic frequency of the glass transition. The maximum (dashed line) is estimated by time-temperature superposition (TTS). If we instead try to reach the glass transition on compression at constant T = 210 K (bottom panel), there is an immediate response to the pressure change but then the α relaxation is practically stuck at a frequency of the maximum loss around 0.05 Hz. On compression (bottom panel), the three last measurements in blue from 10\textsuperscript{-2} Hz take 1 h each and show practically no movement of the maximum loss peak. We interpret this as an indication that pressure is not properly transmitted onto the sample. A likely scenario is that the sample or the pressure-transmitting liquid simply becomes too viscous to transmit pressure efficiently. This aspect is critical for viscous liquids, in which the structural relaxation is extremely dependent on temperature and pressure, that is, the α relaxation time often changes with an order of magnitude as a response to small changes in temperature and/or pressure.

The fact that pressure is sometimes not transmitted properly onto the sample volume when entering the glass on compression is also illustrated by the NS measurements. Neutron TOF measurements from IN6 are presented in Fig. 6, and were carried out simultaneously with the dielectric spectra in Fig. 5. In Fig. 6, we compare the dynamic structure factor summed over Q, S(ω), of DPG at 210 K and 200 MPa coming into the glass on compression or on cooling. The spectra have
FIG. 5. Double logarithmic plots of the imaginary part of the capacitance as a function of frequency for DPG when approaching the glass transition either on cooling at constant pressure (top) or on compressing at constant temperature (bottom). Both graphs illustrate the sample relaxation towards equilibrium. Dashed line in the top panel at 210 K and 200 MPa show the maximum of the loss peak located at a frequency corresponding to $T_g$ and is obtained through time-temperature superposition (TTS). In the bottom panel, the dashed line corresponds to the expected dielectric spectrum for the equilibrated liquid at the glass transition state point; however, this state point is never reached.

been grouped for constant values of $Q$ in the range 1.2-2 Å$^{-1}$. Since the same behavior is observed for all spectra for each value of $Q$, we decided to sum over $Q$ in the data shown in Fig. 6 to improve statistics. The spectrum on compression after a waiting time of more than 5 h does not match the curve shown by the equilibrated glass formed on cooling, which is in agreement with the simultaneous dielectric data shown in Fig. 5. Unlike the long acquisition periods required to obtain well-resolved TOF spectra (up to a hour of hours, depending on the neutron flux), full dielectric curves from 100 mHz to 1 MHz can be measured in 10 min approximately. Dielectric spectroscopy offers quick and precise information about whether or not the sample is in thermodynamic equilibrium and represents a significant advantage compared to standard neutron techniques. These results indicate that without the knowledge from simultaneous DS one might erroneously interpret the changes in $S(\omega)$ being due to a path dependence in the $(P, T)$ phase diagram.

The dielectric signal from the combined cell also provides fast information if a sample is prone to crystallization and also if mixing of the pressure liquid and sample occurs. In case the sample crystallizes, we will immediately observe a reduction of the loss peak, while the presence of the electrical conducting signal or unexpected broadening of the relaxation curve might indicate the mixing of the sample with pressure-transmitting liquid, alerting about the loss of tightness of the floating barrier. The HP DS-NS cell also provides information immediately if there is a pressure leak, in particular when the plug is subjected to several cycles up to 400 MPa. Leakage in the plug is presumably induced by formation of cracks on the sealing epoxy resin used for feedthroughs. In any case, the failure of the plug is certainly unpredictable since several factors could contribute to the loss of tightness.

As we mentioned above, another practical benefit of monitoring the capacitance of the sample is being able to rapidly detect possible contamination of the sample reservoir with pressure-transmitting liquid. In one of the measurements on DPG at 400 MPa, we detected a strange and very strong conductivity signal entering the dielectric spectrum at low frequencies. We concluded that Fluorinert, the pressure-transmitting liquid, was entering the sample compartment due to loss of floating barrier tightness, an example of a scenario which is difficult to detect with neutrons alone.
IV. CONCLUSIONS

We have designed, constructed, and tested an experimental setup to perform simultaneous neutron and dielectric spectroscopy experiments at high pressure. The performance of the cell has been successfully proved in a broad temperature range and up to a maximum pressure of 500 MPa. It opens new possibilities to address studies on dynamic processes in an extremely broad range of temperature and pressure changes. Moreover, measuring the dielectric signal of the sample is an excellent internal probe to get insight into the state of thermodynamic equilibrium after temperature and pressure changes.

As an application to discovery of new physics, we have successfully employed this setup to explore the phase diagram of simple van de Waals liquids in an unprecedented range of temperature and pressure scales. Our results agreed with the predictions of the isomorph theory and allowed reduction of the phase diagram to a single variable, in which fast and slow relaxations are controlled by a single parameter.

The cell will be available through the ILL user program. For future collaborations or getting information about the availability of the setup, please contact Henriette Wase Hansen or Kristine Niss.

ACKNOWLEDGMENTS

This work has been funded by the Danish Council for Independent Research (Sapere Aude: Starting Grant). The authors gratefully acknowledge use of the facilities at the Institut Laue-Langevin (Grenoble, France) through the Long Term Proposal LTP-6-7. The authors would like to thank the support from Jean-Paul Gonzales and Claude Payre (Service for Advanced Neutron Environment at the ILL) and Sébastien Vial (Diffraction Group at the ILL). We also acknowledge the significant support of the Instrument Control Service (SCI) at the ILL. We thank Preben Olsen, Ebbe H. Larsen, and Torben Rasmussen (workshop at IMFUFA, Department of Science and Environment, Roskilde University) for their support on the design and manufacture of the high-pressure cell. Christiane Alba-Simionesco is also acknowledged for fruitful discussions.

14 M. Béé, Quasielastic Neutron Scattering (Adam Hilger, United Kingdom, 1988).
20 See http://www.ill.eu/instruments-support/instruments-groups/instruments/info for more details on how the spectrometer IN6 works.
22 See https://www.ill.eu/instruments-support/instruments-groups/instruments/in16b/description/instrument-layout/ for more details on how the spectrometer IN16B works.
36 See https://www.ill.eu/fr/users/call-for-proposals/ for information about the different options for submitting a proposal to the ILL.