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Quenching device for electrolytic aqueous solutions

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Abstract. A simple device based on splat cooling to liquid nitrogen temperatures is presented. Its application to the amorphization of binary aqueous solutions by fast cooling is demonstrated. The fraction of amorphous material obtained is 90% in eutectic solutions. Diffraction patterns of the vitrified solutions are presented and discussed.

1 Introduction

1.1 Amorphizing solutions

Glassy solutions of salts in water have been studied for decades [1–5]. Many highly hygroscopic salts typically show a glass forming region [5]. In this region they typically show a good ability to form glasses when cooled. This is the case for LiCl-water solutions with concentration close to the eutectic [1,6]. The composition/temperature phase diagram of solutions, be they metal alloys or salts in water, usually exhibits a eutectic point (E), which is the point of lowest temperature at which the liquid is stable. If crystalline nucleation is slow enough, it is possible to super-cool the liquid below this point. For aqueous LiCl solutions this can be achieved by cooling bulk quantities even at moderate rates of 20 K per hour. Thereby a glassy state can be obtained. In reference [1] the change in glass transition temperature (T_g) with concentration is compared for a number of different solutions, and they attempt to explain the difference in the behaviour of these solutions. We are not aware of a formula for the relation between T_g and concentration that gives satisfactory results for the solutions studied here. However, formulas have been proposed for glass forming polymer solutions [7,8].

Nevertheless, the NaCl-water system could not be amorphized so far, as slow cooling leads to the crystallisation of either pure ice, salt or dihydrate. There is a claim

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by [9] that concentrated aqueous solutions, including those of sodium chloride, have been amorphized by cooling rates as low as 60 K s^{-1} . However, amorphization was not determined by diffraction, but by visual inspection of the transparency of the cooled droplet, by the method described in [10]. This is known to give inaccurate results when the sample is micro-crystalline [11]. Transparent samples are expected to be liquid or amorphous. However, this is known to give inaccurate results since the sample may appear transparent even though micro-crystals are present, as observed in diffraction experiments. If the size of crystallites is much smaller than the wavelength of visible light or if the indices of refraction of the amorphous and crystalline components are similar enough, then the drops may remain clear although diffraction experiments reveal the existence of a crystalline part [11].

The difficulty to obtain amorphous NaCl-water solutions, although striking compared to other solutions, seems more understandable given that pure water is even more difficult to amorphize (see Sect. 1.3). There has been an ongoing debate on the relation between the glass transition temperature, distinct phases of amorphous water and the structure of the deep supercooled state [12–14]. Water is a common substance with properties that seem unusual when compared to other pure substances [15]. In the following, we discuss the occurrence of amorphous ice and solutions, and how they can be produced in laboratory experiments.

1.2 Occurence of amorphous water in nature

Pure water is a "bad glass former", i.e. extreme cooling rates are required to amorphize it, and various techniques have been proposed to achieve this. In the next subsection we will briefly present the techniques which work for pure water and discuss whether they are suited for solutions.

Amorphous ice is likely the most common phase of water in interstellar space [16]. In the solar system, the total mass of all types of ices is expected to be about twice that of the rocky material [17]. Recent work has suggested that ice is expected to play a crucial role in the accretion process leading up to the formation of planet sized bodies [18-20]. It has been demonstrated in a numerical model that condensation of vapour is an efficient growth mode towards the formation of decimeter sized pebbles [21]. Thus crystalline and amorphous ices occur in heterogeneous extreme environments ranging from small particles to planetary bodies in protoplanetary disks surrounding young stars. The interaction of solid phases of water with other elements in these conditions ought be taken into account. Salts are an excellent probe to test this interaction, because they are naturally abundant and soluble in liquid water. The question whether ices can include ions ought to be addressed.

Water crystallizes into ice I_h when cooled below the freezing point at ambient pressure. Freezing involves formation of a critical crystal nucleus (nucleation) and its growth. The nucleus may be formed by homogeneous or heterogeneous nucleation. Homogeneous nucleation is the formation of nuclei without preferential nucleation sites. It occurs spontaneously at $T_{\rm hom} = 235 \,\mathrm{K} = -38$ °C at 1 atm. If minor impurities are present, heterogeneous nucleation occurs on their surface and water freezes at higher temperatures [22]. Since water cannot be supercooled below the homogeneous nucleation amorphous material.

1.3 Techniques to obtain amorphous pure water and solutions

The first technique used to amorphize pure water was vapour deposition which leads to the formation of a low density amorphous form called amorphous solid water. This phase is similar to the hyper-quenched glassy water obtained upon hyper-quenching liquid droplets with cooling rates larger than 10^5 K/s [23]. Together these phases are called low density amorphous water (LDA). In this phase, the water molecules mostly retain their tetrahedral coordination [24]. This process is expected to occur in the mesosphere (≈ 50 to 100 km altitude), the coldest layer of the atmosphere, where temperatures can drop below 130 K [22,25]. It is also common in outer space, where water is often observed in an amorphous form [16].

Pure water can also be amorphized by compression (to $\approx 1 \text{ GPa}$) of ice I_h at low temperature [26,27]. The unannealed phase thus obtained is a strained and unrelaxed high density amorphous (uHDA) form of solid water [14]. This can be relaxed by annealing to $\approx 125 \text{ K}$ under pressure yielding the expanded HDA (eHDA) form. Upon annealing to about 115 K at ambient pressure, uHDA transforms to LDA [28,29].

Nevertheless, in the case of viscous solutions these two techniques encounter difficulties. The first one, vapour deposition or epitaxy, is a rather slow technique in terms of the amount of sample produced and it cannot be used easily to form highly concentrated amorphous solutions of salts in water at a given concentration. The second one is unsuited because ice I_h expels salts upon crystallisation, thus a homogeneous crystalline precursor at ambient pressure cannot be produced [30]. Once phase separation of a solution reaches a macroscopic scale, for instance by crystallisation of ice I_h from the solution, mixing times become very long compared to the time scale of most experiments, because the diffusion of ions in the glass is very slow at low temperatures. Hence this technique is not suited to produce a glass with uniform distribution of ions.

Two techniques which are suitable to amorphize solutions are jet freezing of emulsions and splat-cooling. It has been shown that pure water and dilute solutions can be amorphized by jet freezing droplets in emulsion [31]. For the emulsion a liquid cryomedium such as n-heptane has to be used. This method has been used for solutions of LiCl [32]. The main inconveniences are the difficulty to separate the sample from the cryomedium and to avoid crystallisation of the sample droplets in the process.

The technique used in this work is similar to the splat-cooling technique, which was originally developed for metals in the 1960s [33]. The group of Mayer at the University of Innsbruck (Austria) were the first to apply this approach to pure water and aqueous solutions [23]. This technique is based on the rapid cooling of droplets (called splat) on a solid substrate. The three possible regimes of splat cooling are ideal cooling, intermediate cooling and Newtonian cooling. The boundaries between these regimes are expressed in terms of the dimensionless Nusselt number, which is Nu = h s/K, where h is the heat transfer coefficient for the interface between splat and substrate, s is the thickness of the splat layer and K is the thermal conductivity of the splat.

In the first case, known as ideal cooling $(Nu \to +\infty)$, the cooling rate is determined completely by the droplet's size and its thermal properties. In reference [33] it has been shown that for perfect contact $(h = +\infty)$ between splat and a semi-infinite substrate, whose density, specific heat and thermal conductivity are temperature independent, the cooling rate (\dot{T}) , apart from constants for material properties, varies as $\dot{T} \propto z^{-2}$, z being the distance from the interface. Hence larger cooling rates are achievable closer to the interface, this means that in thinner splat the minimum cooling rate will be larger.

Newtonian cooling is the opposite limiting case $Nu \ll 1$, where the temperature gradient in the droplet is negligible and the cooling rate is determined entirely by the heat flow at the interface. Jones [33] refers to the region in between, where Nu is of the order of unity, as intermediate cooling. The Newtonian and ideal regime are illustrated in Figure 1. According to finite difference calculations by [34] for iron freezing on copper, Newtonian cooling prevails for Nu < 0.015, and ideal cooling for



Fig. 1. The Newtonian and ideal cooling regimes of a droplet (splat) on a substrate during splat cooling.

Nu > 30. These calculations also show that for Newtonian cooling $\dot{T} \propto z^{-\alpha}$, where $\alpha \ge 1$ is a constant.

2 Quenching procedure

2.1 Experimental set-up

Our device for fast quenching is depicted in Figure 2 and a schematic identifying the parts is given in Figure 3. It exploits the rapid cooling of micrometer sized droplets on a pre-cooled solid substrate. The surface, on which droplets are projected, is a circular plate of an aluminium-copper alloy. This alloy was chosen for its high thermal conductivity ($K_{\rm Du} \approx 100 \text{ W/m/K}$ at 80 K) to which the cooling rate of the sample is proportional [33]. The plate is placed inside a cylindrical vacuum chamber (diameter 330 mm, height 200 mm). The pressure inside is of the order of 1 mbar. The chamber is cooled by immersing its lower part in a liquid nitrogen bath. Cooling the device from room temperature (300 K) to 80 K takes about 1.5 hours. Moreover, the sample may be pre-cooled to 0 °C. This lowers the vapour pressure of the solution, thereby reducing evaporation in the vacuum and variations of the final concentration in the droplets.

The quenching proceeds in two steps: injection onto the plate and collection of the amorphous sample. Upon activation of the pneumatic valve, the sample solution is injected through a nozzle onto the cooled aluminium-copper plate. Commercial nozzles produced by Fluidics were used in our experiment. This type of nozzles is widely used for fuel burners and presents the advantage that for a given fluid (viscosity) and intake pressure, the flow rate has been calibrated. Also, the nozzles are available for a large range of viscosities, hence the quenching device can easily be adapted to the fluid to be splat-quenched. The flow rate through the nozzle at 80 bar is about 2.51/h, and the velocity of the droplets can be estimated as $v = \sqrt{2p/\rho} \approx 120 \text{ m/s}$. The cone of injection, depicted in Figure 3 by two lines surrounding the droplet, has an opening angle of 80°. A serious problem is corrosion, despite the fact that all materials in contact with the salty solutions are made of stainless steel. The device has to be cleaned after each use. To prevent corrosion the entire device was purged with pure water at least 4 times. The nozzle was removed, disassembled, and its parts rinsed in an ultrasonic bath in pure water, which was changed several times.



Fig. 2. Photography of the quenching device atop the container serving as liquid nitrogen bath. The parts are identified by numbers as: 1: vacuum chamber, 2: cryoplate, 3: nozzle, 4: valve, 5: actuator, 6: control unit, 7: manometer, 8: nitrogen funnel.

The pneumatic actuator is controlled by an electronic interface via an electronic actuator produced by Topindustrie (France). The system allows the operator to choose an injection time with a precision of 0.1 s.

In order to collect the amorphous sample we proceeded carefully to guarantee that the sample remains close to liquid nitrogen temperature. The vacuum was broken by flooding liquid nitrogen into the chamber through a pipe connected to a funnel. This lead at first to a turbulent flow of solid nitrogen in the chamber. Small patches of sample were torn off the plate and formed a plaque, whose temperature may have risen higher than that of the powder sticking to the plate. This part was discarded. The rest of the sample was slowly immersed and remained at liquid nitrogen temperature. This part of the sample was reduced to fine powder, recovered under N_2 vapour, collected in plastic containers filled with liquid nitrogen, and stored in dewars. Roughly 1 g of powder was produced in each quenching procedure.

Tests were performed varying four parameters in order to determine the conditions that allow to amorphise most of the sample for a given concentration and composition. These parameters are the injection pressure, the injection time, the pre-cooling temperature of the sample, and the surface roughness of the plate.

In our set-up, a pressure of up to 200 bar of helium gas was applied to the solution in the tube to drive it through the nozzle. Calibration experiments were performed to measure droplet size and amorphisation degree at different pressures. The droplet size was measured with an optical microscope on samples injected directly into liquid nitrogen. The result is shown in Figure 4, which shows that the droplet size decreases with applied pressure. The diameter of the droplets produced at 70 bar was measured to lie between 10 and 30 μ m (Fig. 4) with an average droplet size of about 20 μ m. The average droplet size was estimated from photographs taken with an optical microscope through a glass plate onto the sample droplets in liquid nitrogen. The



Fig. 3. Schematic of the quenching device. The sample solution in the tube above the nozzle is pre-cooled to about 0 °C. The nozzle projects small droplets of diameter ≈ 10 to $30 \,\mu\text{m}$ onto the aluminium-copper plate at 80 K, with a spraying angle $\theta \approx 80^{\circ}$.



Fig. 4. Left: droplet size as a function of injection pressure for pure water (blue circles) and NaCl solution with R = 10.5 (red diamonds). The molar ratio is defined in Section 2.2. The droplet size was measured on samples injected directly into liquid nitrogen. The width of the distribution of droplet sizes is about 5 or $10 \,\mu$ m. Right: photograph of the droplets produced at about 40 bar seen through an optical microscope.

spread of the distribution of droplet sizes at a given pressure is 5 to 10 μ m. Therefore it is reasonable to approximate the droplet size as about 20 μ m at 80 bar.

The injection time t was also varied. Times from 0.5 to 3.0 s were tested. Since the amount of material injected is roughly proportional to the injection time, the latter controls also the thickness of the layer of splat. Thicker layers would cool more slowly and thus nucleation of ice can occur. Thinner layers yield less sample and are more difficult to collect. The injection time was chosen as a compromise between sample amount and cooling speed.

The largest fraction of amorphous material of NaCl·10.2D₂O was obtained in conditions of 80 bar with 0.6 s injection time, on a surface roughened with sandpaper (grade P400, average particle diameter of 35 μ m). For a droplet diameter of about 20 μ m, the Nusselt number is \approx 1, hence we are in the intermediate regime of cooling. Using the data of [34] and [33], we estimate our cooling rate to be approximately 10^4 to 10^5 K s⁻¹. Using another approach where $\dot{T} = \Delta T/\tau$, where the temperature difference for the droplet is $\Delta T \approx 200$ K, the characteristic cooling time is $\tau = s^2/D$, s being the drop size and $D = 1.2 \times 10^{-7}$ m²/s the thermal diffusivity of water at 25 °C, we obtain $\dot{T} \approx 6 \times 10^4$ K/s which is consistent with the previous estimate.

2.2 Preparation of the solution and powder

Solutions of NaCl were prepared by dissolving salt in light or heavy water. For most X-ray diffraction experiments we used solutions in H₂O. D₂O solutions were used for neutron scattering, because of the large incoherent scattering of H. The D₂O was at least 99.75% pure and H₂O was filtered by a Millipore DirectQ. Its resistivity is greater than 18.2 M Ω ·cm at 23 °C (Grade 1 of the ISO 3696 standard). The concentrations are discussed in terms of the molar ratio R = n(water)/n(salt).

For sodium chloride (rock salt) the stable eutectic concentration in H₂O is R = 10.7 close to the maximum solubility which is at R = 9.05 at ambient temperature and atmospheric pressure. In D₂O the solubility limit is 9.7, our experiments indicate that the metastable eutectic is at 10.2. Solutions with R in the range of 9.05 to 11 were produced to determine for which concentration the best amorphisation could be achieved. To ensure complete and homogeneous dissolution, all solutions were plunged for about ten minutes in an ultrasonic bath.

3 Amorphous aqueous solutions

The structure of amorphous samples obtained using the procedure described in the last section was studied by X-ray and neutron diffraction at low pressure (p < 1 atm). First, we briefly describe the diffraction experiment. Second, we give an estimate of the fraction of amorphous material.

3.1 Diffraction on amorphous samples

In order to check the degree of amorphization, the samples were studied at ambient pressure by X-ray diffraction with a commercial X-PERT powder X-ray diffractometer at IMPMC, equipped with a low-temperature Anton Paar chamber. For samples intended for neutron diffraction experiments, a small amount of quenched solution was systematically recovered and measured as above by XRD. The sample was placed on a sample holder made of nickel plated copper, which was cooled to ≈ 80 K. During transfer the temperature of the sample did not exceed 95 K. This has been tested



Fig. 5. X-ray diffraction patterns of quenched NaCl·10.2D₂O and NaCl·9.5H₂O at 80 K in the cryochamber. The samples were precooled to $\sim 0^{\circ}$ C. They are the ones with the largest fraction of amorphous material.

using a mixture of iso-pentane and n-pentane which undergoes a glass transition for $T \approx 100 \,\mathrm{K}$. The diffractometer uses Cu $K\alpha$ radiation with an incident wavelength of $\lambda = 1.5418$ Å, set up in $\theta/2\theta$ geometry. The X-ray diffraction patterns of NaCl solution in H₂O and D₂O are shown in Figure 5.

The neutron study was carried out on the high-intensity two-axis neutron diffractometer D20 [35] at the high-flux reactor of the Institut Laue-Langevin (ILL, Grenoble, France). The set-up is the same as that described in [36]. The monochromator was set to a wavelength of 1.87 Å with average resolution of $\delta d/d = 3 \cdot 10^{-3}$. A helium flow cryostat, pre-cooled at 80 K with a pressure of approximately 100 mbar of helium exchange gas, was used as sample environment. The powder sample was loaded into a cylindrical vanadium cell of 6 mm diameter kept under liquid nitrogen. Three solutions of different concentrations close to the eutectic one were measured in this experiment. The diffraction data were normalised using the empty vanadium cell, empty cryostat, and vanadium standard measurements. In order to remove the background contribution from the cryostat and correct for detector efficiency, the data were reduced using the formula given in [36]. The neutron diffraction patterns of NaCl solution in D₂O and of pure D₂O ice I_h are shown in Figure 6.

In order to identify the phases present in the samples, the diffraction patterns of the known crystalline phases of the system at ambient pressure were calculated with FullProf [37,38]. The structure data for ice I_h (hexagonal, space group $P6_3/mmc$) was taken from [39]. That of NaCl is face-centered cubic (space group $Fm\bar{3}m$). The sodium chloride dihydrate structure (monoclinic, "probable" space group $P2_1/c$) of [40] was used.



Fig. 6. Neutron diffraction patterns of quenched NaCl·R D₂O and pure ice I_h (top) at 80 K measured on D20 at ILL. The composition is indicated on the right. During the measurement the pressure is that of the helium exchange gas (≈ 100 mbar). The sample with R = 10.2 contains more amorphous material with respect to the crystalline I_h. Hydrate peaks appear in the pattern of the sample with R = 9.7 but not in that with R = 10.2. The first four clearly visible hydrate peaks are marked by asterisks.

3.2 Fraction of amorphous material

Figure 5 shows the X-ray diffraction patterns of samples which show the highest fraction of amorphous versus crystalline material. The first samples produced were obtained from NaCl-H₂O solutions with compositions R between 9 and 11. This concentration range was chosen to bracket the stable eutectic concentration (R = 10.7). The samples with 9.5 $\leq R \leq$ 9.8 contained the largest fraction of amorphous material.

For D₂O the solubility limit of NaCl is R = 9.7 at room temperature and ambient pressure. Trials at different concentrations showed that the largest amorphous fraction could be obtained for R = 10.2. This was confirmed in our experiment on



Fig. 7. Neutron diffraction pattern of NaCl-10.2D₂O measured on D20 at ILL [36] at 100 mbar (red points). The sum of the fits of amorphous and crystalline contributions is shown in blue on top of the data. The crystalline part is estimated to be about 8% of the sample material according to the fit of thirteen ice peaks with pseudo-Voigt profile functions, shown in green below the data.

 D_2O at ILL. Figure 6 shows the neutron diffraction patterns obtained for pure hexagonal ice of D_2O , for NaCl·9.7D₂O and for NaCl·10.2D₂O. Although both salty samples contain amorphous material, a larger fraction is present in the sample with R = 10.2. The sample with R = 9.7 shows larger ice I_h peaks and also NaCl dihydrate peaks. The latter observation is coherent with the phase diagram of NaCl-H₂O [41,42]. The fact that R = 10.2 solutions amorphise without forming dihydrate indicates that the rate of formation of the dihydrate at this concentration is slower than the quenching rate of our device.

From the above observations we conclude that the isotopic effect of substituting D for H in these solutions shifts the composition of best amorphisation from R = 9.5 in H₂O to R = 10.2 in D₂O.

Figure 7 shows the neutron diffraction data of NaCl·10.2D₂O at 90 K and a fit to the data. The data reveal both peaks of crystalline material and amorphous features of the solution. By comparison with FullProf simulations of ice I_h peaks, we conclude that all the peaks correspond to ice I_h , although with some texture. Therefore thirteen of the ice peaks in the neutron data at 90 K were fitted individually with pseudo-Voigt profile functions using the least squares algorithm of GNUPLOT 4.6 [43]. The respective intensities were obtained by integrating the fit of the ice peaks and the structure factor of amorphous material. This gave amorphous and crystalline volume fractions of 92% and 8% respectively. The texture observed in the sample may introduce an error of few percent points to the estimate of the fraction of crystalline material in the sample. We estimate that the error on the fraction above is approximately 2%. Hence at most 10% are expected to be crystalline, and at least 90% amorphous.

The neutron data provide evidence that the crystalline part forms during the hyper-quenching process since the measured ice peaks correspond to deuterated ice I_h . This cannot be formed by condensation of water from a humid ambient atmosphere during loading of the sample, which would be H_2O ice. This deuterated ice must have formed from the injected sample. It is possible that this part of the sample heated up when it detached from the plate when the vacuum was broken. Another explanation would be that the layer of sample formed after injection is not of uniform height. In the areas where the sample layer is thicker cooling may be slower thus allowing the nucleation of pure ice.

A few samples of pure water quenched with our device have been measured by XRD. All of them were highly crystalline. This is consistent with the cooling rate estimated to be $\approx 10^4 \,\mathrm{K \ s^{-1}}$ (see Sect. 2.1) which is lower than the minimal cooling rate required to amorphise pure water ($\geq 10^5 \,\mathrm{K \ s^{-1}}$).

We have found that concentrated NaCl solutions amorphise at ambient pressure in a high density structure similar to the HDA form of pure water [36,44], in agreement with similar observations for LiBr and LiCl solutions [45,46]. The samples produced with this device have been used in high pressure neutron diffraction studies [36,47].

4 Conclusion

The quenching technique described here allows to amorphise highly concentrated viscous solutions of NaCl in water with an estimated quench rate of 10^4 to 10^5 K s⁻¹. The best amorphous samples were obtained for R = 9.5 in H₂O and 10.2 in D₂O. An amorphous content of 92% was obtained in D₂O.

This method may be applied to obtain amorphous forms of other aqueous solutions or solutions with viscosity and surface tension comparable to that of the NaCl-water solutions. We believe that the residual crystalline impurities are due to a small amount of large droplets which suffer slow cooling times, or a bad contact of some part of the projected sample to the cold-plate. With further investigation this device might be improved and applied to produce amorphous forms of other aqueous solutions which could not be produced so far in large quantities.

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