Carbon-nanotube based materials for protein crystallisation

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ABSTRACT

We report on the first use of carbon-nanotube based films to produce crystals of proteins. The crystals nucleate on the surface of the film. The difficulty of crystallising proteins is a major bottleneck in the determination of the structure and function of biological molecules. The crystallisation of two model proteins and two medically relevant proteins was studied. Quantitative data on the crystallisation times of the model protein lysozyme are also presented. Two types of the nanotube film, one made with the surfactant Triton X-100 (TX-100) and one with gelatin, were tested. Both induce nucleation of the crystal phase at supersaturations at which the protein solution would otherwise remain clear, however the gelatin-based film induced nucleation down to much lower supersaturations for the two model proteins with which it was used. It appears that the interactions of gelatin with the protein molecules are particularly favourable to nucleation. Crystals of the C1 domain of the human cardiac myosin-binding protein-C that diffracted to a resolution of 1.6Å, were obtained on the TX-100 film. This is far superior to the best crystals obtained using standard techniques, which only diffracted to 3.0 Å. Thus, both our nanotube-based films are very promising candidates for future work on crystallising difficult-to-crystallise target proteins.
INTRODUCTION

We report on the first use of carbon-nanotube based materials to form crystals of biological macromolecules. Crystallising these large molecules, particularly proteins, is a key problem in modern biology. Protein structures are pivotal to the success of rational drug design and to other biotechnology applications, and so international structural genomics/proteomics projects have set out to determine the structures of all the proteins in our genome. The most powerful technique for determining protein structure is X-ray crystallography. This of course requires high quality crystals of the proteins. Producing such crystals is a very difficult task that has long been a major bottleneck to progress in structural biology. In the era of genomics/proteomics, this problem is acute: only about one in five of the proteins obtained as pure solutions have been crystallised.\(^1\) Thus, there is an urgent requirement for new ideas and tools. Here we show that a carbon-nanotube based material with nanometre sized pores is an effective and versatile tool for the crystallisation of proteins.

Crystallisation is a first-order phase transition and so proceeds via nucleation of the crystal, followed by its growth. Without nucleation no crystallisation can occur, thus it is important to be able to induce nucleation, and we also need to control the amount of nucleation.\(^2,3\) This is because a common problem in crystal growth is the formation of excess nuclei, which leads to the production of large numbers of small, useless (for diffraction studies), crystals instead of the desired few large ones.\(^1\) Nucleation has only really been systematically for one protein, lysozyme; this is discussed in a recent review by Sear\(^2\). For this protein it is clear that nucleation is always or almost always heterogeneous, \textit{i.e.}, the crystal forms in contact with a surface\(^2\). The situation is less clear for other proteins, although there is no reason to expect lysozyme to be exceptional. This finding of nucleation on surfaces has led to the idea of deliberately adding substances to a solution in order to provide surfaces where nucleation readily occurs. Such substances are called nucleants.
The ideal nucleant for protein crystallisation should have the following properties:\(^4\)

1) It should act as a nucleant for *many* proteins, not just one. As proteins are diverse, this is a demanding requirement.

2) It should allow control over the number of nuclei that form, thereby promoting the growth of only one or a few crystals.

3) The supersaturations at which nucleation occurs on the nucleant should be amenable to control, in order to nucleate crystals under conditions that are as close as possible to ideal growth conditions.

Starting with the work of McPherson and Schlichta,\(^5\) many diverse materials have been used as nucleants for protein crystallisation,\(^6,7,8,9,10\) even including nanoparticles\(^11\), and some success has been achieved. However, the ideal nucleant has not yet been found. Our aim was to select a novel nucleant based on knowledge of the microscopic details of nucleation. To do this, we considered the results of computer simulations. These showed that pores of size comparable to that of the nucleus\(^12\) promote rapid nucleation. This is a generic effect that should apply to all proteins, as well as to other molecules. Proteins are a few nanometres across, and the nucleus of a protein crystal phase is expected to be a few protein molecules across.\(^2\) Thus a surface with pores \(\sim 10\) nm across are good candidates to induce the nucleation of protein crystals.

In earlier work, we used two types of media with nanoscale pores as nucleants: etched silicon\(^13\) and bio-glass.\(^4\) In both cases the surfaces are composed of pores of typical size a few nanometres across but each pore has a different size and shape. We found that bio-glass induced crystallisation of the largest number of proteins ever crystallised using a single nucleant.\(^4\) However, the pore sizes and surface chemistry of this particular medium are not easily controlled.

So, we sought a material with a surface with pores whose size and surface chemistry could be controlled. We found one: films of entangled carbon nanotubes.\(^14,15\) These films are currently being investigated for a number of applications\(^16\) due to their, for example, mechanical\(^17\) and electrical\(^18\)
properties. We refer to these films as buckypaper. Buckypaper is a disordered mat of entangled, and typically coated, carbon nanotubes. See Fig. 1. Carbon nanotubes have diameters of a few nanometres but lengths of hundreds of nanometres. Here, we present results first of the characterisation of the buckypapers and then of their use as nucleants.

**MATERIALS AND METHODS**

**Synthesis of opaque buckypaper with Triton X-100** (used for all except the results in Fig. 6) Single-walled carbon nanotubes (SWNTs), synthesized via high-pressure CO conversion (HiPCO), are obtained from Carbon Nanotechnologies Inc. (CNI). The buckypapers were made from the suspension of 0.04 mg/ml of SWNTs and 0.4 mg/ml of Triton X-100 with deionised water in a total volume of 500 ml. The TX-100 is first diluted with 40 ml of deionised water. The prepared SWNTs were then mixed with the TX-100 solution and sonicated by probe sonicator at 28-30 W for one of two different times: 4 minutes \((t = 4 \text{ min.})\) and 40 minutes \((t = 40 \text{ min.})\). Each prepared suspension was then diluted with deionised water up to 500 ml and sonicated in a batch sonicator for 30 minutes. The suspension was then ready to make a buckypaper. The suspension was filtered through a 0.22 µm filter membrane under high pressure from a water pump. The filter was obtained from Millipore (a nylon filter membrane) with a diameter of 47 mm. When the entire amount of suspension was passed through the filter, 2 liters of deionised water was then passed through it. The dried buckypaper was then kept in the oven at 65-70 °C for 12 hr. Finally, the buckypaper was removed from the membrane by peeling it off and is then ready.

Passing deionised water through the filter removes the more labile TX-100. Note that substantial amounts of TX-100 remain to coat the SWNTs after rinsing of both preparations. The amount remaining of TX-100 was confirmed by thermogravimetric analysis (TGA) in air. We find that the buckypaper is approximately 16 % TX-100 by mass.

Annealing removes the surfactant. The buckypapers, made from suspensions with a probe sonication time of 40 minutes, were annealed under an argon atmosphere with a heating rate of 5 °C/min and held at 600 °C for 90 minutes.

**Transparent buckypaper with TX-100 (200-nm thick)** The suspension was prepared from a dilution of 1 ml of suspension from probe sonication time of 40 minutes \((t = 40 \text{ min.})\) with deionised water up to 100 ml and then was sonicated with the probe sonicator at 28-30 W for 30 minutes and with
a batch sonicator for 1 hour. The suspension was then filtered through a 0.22 µm filter membrane. When
the entire amount of suspension was passed through the filter, 500ml of deionised water was then
passed through it. The dried buckypaper was then kept in the oven at 65-70 0C for 12 hr. To remove the
buckypaper from the membrane, the nylon membrane was dissolved in acetone. The free-standing
buckypaper was attached to a cover glass.

**Gelatin buckypaper** The buckypapers were made from a suspension of SWNTs and gelatin
(Sigma, porcine skin, type A) in deionised water. We started with 1 g of gelatin, which was diluted with
40 ml of deionised water and then autoclaved at a temperature of 121 0C for 15 min. at a pressure of 15
psi (100kPa). The autoclaved gelatin suspension was then centrifuged twice at 3000g, each time for 40
minutes. We then took the supernatant of this centrifuged suspension and added 0.01g of HiPCO
SWNTs. The resulting dispersion was sonicated at 50 W for 40 minutes by probe sonication. The
sonicated dispersion was then diluted with deionised water up to 1 liter and sonicated with the batch
sonicator for 1 hour. The suspension was filtered through a 0.22 µm filter membrane. When the entire
amount of suspension had passed through the filter, the dried buckypaper was then kept in the oven at
35 0C for 24 hr. The gelatin buckypaper was removed from the membrane by peeling it off and is then
ready.

We carried out nitrogen adsorption-desorption isotherms of Fig. 2 at 77 K with a calibrated
Coulter SA 3100 Series instrument. Standards were run before and after each buckypaper isotherm.
Judging from the differences between the before and after standard isotherms we estimate an accuracy
of around 2 cm³/g in the volume adsorbed, \( V_{\text{ads}} \), for the isotherms of our samples of mass close to 0.2 g.
The BET surface area and pore size distribution of Fig. 1 were obtained from these isotherms. SEM
(Hitachi S4000) and AFM (NT-MDT) were both performed at the University of Surrey.

**Characterisation of gelatin solutions** Gelatin solutions were prepared as described above. To
characterize the gelatin in the solution we started with a 0.1% gelatin suspension; 1g of gelatin in 1 l of
deonised water, prepared from an autoclaved gelatin suspension as above. This suspension was stirred,
by a magnetic stirring bar, for 15 min at the temperature of 70 0C. Then, 1ml of the warm suspension
was diluted with deionised water up to 1 l. The 1 l suspension was stirred by a magnetic stirring bar for
a further 15 min at the temperature of 70 0C. Then, 10 µl of the suspension was dropped onto a cover
glass (13-mm diameter, Agar Scientific), which was spun at 4000 rpm for 1 min. The surface was
imaged with an AFM. A typical image is shown in Fig. 4.
Crystallisation trials All crystallisation trials were performed using the vapour-diffusion hanging-drop method in EasyXtal Tool (QIAGEN) plates, except for the trials with transparent buckypaper. These trials used standard silanised microscope cover glasses instead of the EasyXtal tool, and vacuum grease for sealing. A 20 mg/ml solution of lysozyme (Sigma/L6876) was prepared by further dilution with deionised water of a 60 mg/ml solution. This solution was centrifuged at 3000g for one hour and filtered through a 0.22 µm Millipore filter.

The crystallisation trials with lysozyme (Fig. 5(A)) and the TX-100 (unannealed and annealed buckypapers) and gelatin buckypapers were all done in the same way, as follows. In the EasyXtal plates each reservoir contained 500 µl of a solution consisting of 0.1 M sodium acetate (Fisher) buffer at pH 4.5, and sodium chloride (Fisher) at various concentrations. For each trial, 1 µl protein solution was mixed with 1 µl of reservoir solution on a lid, which was then inverted and sealed over the reservoir. Nucleants were directly inserted into the droplets. In one experiment, 5 droplets were prepared at each salt concentration, and of these 5, 3 contained the buckypaper nucleant and 2 did not (control droplets). The time to nucleate in this experiment, \( S_i \), in the presence of the buckypaper was then taken be the average of the times that crystals were first observed in the 3 droplets – rare instances where in one droplet either no nucleation occurred or it occurred very rapidly were not counted. The points in Fig. 5(A) are then the mean of 5 repeats of this experiment, i.e., the mean time is \( (S_1 + S_2 + S_3 + S_4 + S_5)/5 \), and the error bars are just the standard deviation of this set of 5 numbers. The control drop curve is obtained in the same way from the results obtained in the 2 control droplets in one set of 5 repeat experiments. The experiments were observed for 2 weeks.

Crystallisation trials with gelatin solution as a nucleant were performed using the EasyXtal plates described above. As above each reservoir contained 500 µl of 0.1 M sodium acetate buffer at pH 4.5 and 3.6% NaCl. Each control droplet was prepared by mixing 2 µl of lysozyme (20 mg/ml) with 2 µl of 3.6% NaCl. These are metastable conditions (see Fig. 5A)). Each sample droplet was prepared in the same way as a control droplet, but with a 0.5 µl droplet of 0.1% gelatin solution added. The experiments were observed for a week.

Trypsin was crystallised from a 30 mg/ml protein solution containing 20 mM Tris pH 8.2. The reservoir solutions consisted of 0.1 M Tris buffer at pH 8.2 and ammonium sulphate. The concentration of ammonium sulphate ranged from 1.08 to 1.32 M. At least four identical experiments were performed for each condition, with and without buckypaper.
**Human myosin-binding protein-C (MyBP-C)** was crystallised from a 10 mg/ml protein solution containing 50 mM NaCl and 20 mM Tris, pH 7.0. The reservoir solutions consisted of 0.1 M HEPES buffer at pH 7.3 and PEG of mean molecular weight 3350. The concentration of PEG 3350 ranged from 15% (w/v) to 20% (w/v) in steps of 1%. Six identical experiments were performed for each condition, with and without TX-100 buckypaper.

Approximately 10 crystals grown on the buckypaper were X-rayed at beam line 10.1, SRS-Daresbury, using an Oxford Cryosystems cryojet at 100K and a MAR 225 CCD detector.

**Non-structural protein 9 of the transmissible gastroenteritis virus (NSP9)** The viral protein was crystallised at 20 °C from a 12mg/ml protein solution containing 200 mM NaCl and 10 mM HEPES, pH 7.5. The reservoir solutions consisted of 0.1M HEPES buffer at pH 7.5, 15% (v/v) isopropanol and PEG 3350. PEG 3350 concentrations were between 16% and 20%, (w/v) in steps of 1%. Six identical experiments were performed for each condition, with and without TX-100 buckypaper.

**RESULTS AND DISCUSSION**

**A. Buckypaper preparation and characterisation** Two types of buckypaper were produced: one with the surfactant Triton X-100 (TX-100), and one with gelatin. Carbon nanotubes are highly insoluble in water thus TX-100 or gelatin are needed to coat the carbon nanotubes in order to render the surfaces hydrophilic and so disperse them in solution. Note that the gelatin buckypaper has more gelatin by mass than carbon nanotubes, and annealing the TX-100 buckypaper to remove the TX-100 significantly increases the contact angle (see Fig. 1(C) and (D)). So, we expect the carbon surfaces in the buckypaper to be coated by TX-100 or gelatin.

Figure 1 shows both an SEM image of a TX-100 buckypaper, and pore-size distributions obtained by Brunauer–Emmett–Teller (BET) analysis of nitrogen adsorption. The nitrogen physisorption isotherm is shown in Fig. 2(A). Note that the pores are the spaces between bundles of coated SWNTs not the cavity
within a single SWNT. From the SEM image of Fig. 1(B) we see that the nanotubes forming the buckypaper are not individual tubes, but bundles approximately 10 nm across. The pore-size distribution can be controlled, within a range, by varying the probe sonication time of the SWNT solution; the pores are on average larger if we only probe sonicate for \( t = 4 \) minutes, than if we sonicate for \( t = 40 \) minutes. After \( t = 40 \) minutes the dominant peak in the distribution of pore sizes is around 9 nm – approximately 3 lysozyme protein molecules across. The BET surface area is also a function of sonication time. It equals 89 m\(^2\)g\(^{-1}\) and 53 m\(^2\)g\(^{-1}\), for \( t = 4 \) and \( t = 40 \) minutes, respectively. Somewhat surprisingly, the area goes down as the pore size goes down. We only used as a nucleant, the TX-100 buckypaper obtained with the longer (\( t = 40 \) minute) sonication time. In order to obtain information on the effect of surface chemistry we also annealed some of this buckypaper in an argon atmosphere at 600 \(^0\)C for 90 minutes. This removed the TX-100 (see Materials and Methods section), thus making the surfaces somewhat hydrophobic (see Fig. 1(D)). It also made the pores smaller on average. After annealing, the dominant peak in the pore-size distribution is at 4 nm, and the specific surface area of the annealed buckypaper is 564 m\(^2\)g\(^{-1}\). Thus, for the buckypaper synthesized with TX-100 we have some control over both the pore-size distribution and the hydrophilicity of the buckypaper, but we cannot control them independently.

Figure 3 shows an AFM height map of the surface of buckypaper made with gelatin. The much larger size of the gelatin molecules, as opposed to the TX-100 surfactant, appears to result in the spaces between the carbon-nanotube bundles being filled in, or at least covered by gelatin. We can get a clear idea of the difference between the TX-100 and gelatin buckypapers by comparing their nitrogen adsorption and desorption isotherms in Fig. 2. We see that the isotherms for TX-100 have a Type IV shape. This shape is characteristic of capillary condensation in mesopores. Mesopores are defined as pores of widths in the range 2 to 50 nm. See Sing et al.\(^{19}\) for the classification scheme for physisorption isotherms. Note that the BET analysis we employ for the TX-100 buckypaper is reasonable for Type IV isotherms (only). For Type IV isotherms the initial adsorption is typically that of layers of nitrogen on
the surfaces of pores, and the large hysteresis loop is characteristic of capillary condensation in pores. Also, note that the physisorption is large, over 100 cm$^3$ per gram of the buckypaper. Finally, if we place a droplet of water on top of TX-100 buckypaper the water can percolate through to the other side. Thus we are confident that the TX-100 buckypaper is a porous medium, which by definition contains a connected network of pores that span the complete thickness of the buckypaper.

The adsorption and desorption isotherms for the gelatin buckypaper are very different. They are close to Type III,\textsuperscript{19} and the total amount of nitrogen physisorbed is an order of magnitude less. Also, if we place a droplet of water on top of gelatin buckypaper, it does not percolate through. Thus we conclude that although the gelatin buckypaper is very rough and contains pores in the sense of deep indentations, see the AFM data in Fig. 3, it is not a porous medium – it does not contain a network of interconnecting pores. It is possible that the large gelatin molecules block the pores. Below we will find that the gelatin buckypaper is a more effective nucleant than the buckypaper made with TX-100, thus we do not require that the material be a porous medium to be an effective nucleant, a surface with roughness/pores of the right lengthscale is enough. Our AFM data, Fig. 3, clearly shows roughness, and hollows, down to lengthscales a little larger than the expected size of the crystal nucleus (the limit of our resolution). Thus we expect that the nucleus of the crystal phase will feel a pore-like concave surface, just as it would in our TX-100 buckypaper. This finding that a true porous medium is not required is consistent with earlier computer simulation work\textsuperscript{12}, which found rapid nucleation in model pores that were simply rectangular cross-section indentations.

Thermogravimetric Analysis (TGA) of buckypaper showed that it contains gelatin and SWNTs in a ratio of approximately 6:4. This, together with solubilisation of the SWNTs by gelatin suggests that the surface of the gelatin buckypaper is largely gelatin. Gelatin is largely the structural protein collagen. We are not aware of any specific interactions between collagen and any of the proteins we study here. However, collagen contains charged amino acids of both signs, as well as hydrophilic and hydrophobic
The triple helices of collagen can also form aggregates of sizes larger than the expected nucleus size of a protein crystal, and with structure on lengthscales from nanometres to hundreds of nanometres. Thus the surface chemistry of the gelatin buckypaper is expected to be complex, and its nanoscale roughness is presumably due to the collagen fibres as well as to the carbon nanotubes. This complex surface chemistry may allow quite strong attractive interactions between the surface of the gelatin buckypaper and the crystallising proteins. The expectation is that attractive interactions will reduce the free-energy barrier to nucleation.

B. Protein crystallisation trials: Lysozyme

Having discussed the surface structure and chemistry of the buckypapers, we present the results of our crystallisation trials, first with the well-studied model protein lysozyme, then with the less well studied but still common protein trypsin, and finally with two difficult-to-crystallise proteins. Figure 5 presents the first quantitative results for the effect of a nucleant on the time to crystallise of a protein. The results are for crystallisation of lysozyme, at 25°C, with 20 mg/ml protein in a 0.1 M sodium acetate buffer at pH 4.5. The time until the first (5–10 μm across) crystal is visible, is plotted as a function of the salt concentration. The crystallisation conditions and protocols are described in the Materials and Methods section. The solubility of lysozyme decreases as the salt concentration increases so at fixed lysozyme concentration, increasing salt concentrations correspond to increasing supersaturations. At 25°C, tetragonal crystals coexist with a solution of approximately 20 mg/ml in the presence of 2% NaCl, thus all the salt concentrations at which we find nucleation are quite deep into the supersaturated regime. For example, at 4% NaCl, the solution at coexistence has a concentration near 5 mg/ml.

Crystallisation is observed at NaCl concentrations down to 3.6% with the gelatin buckypaper and down to 4.4% for the TX-100 buckypaper. Both are lower than the 4.8% salt concentration which is the lowest at which we find crystallisation in our control experiments, without the buckypaper nucleant. Thus we conclude that our buckypaper nucleants can induce crystallisation at low supersaturations, at
which no nucleation would occur in their absence. In addition, the buckypaper produced with gelatin is significantly more effective than that with the surfactant TX-100. At conditions such that nucleation occurred both with and without a nucleant, the time to observe a (5–10 μm) crystal was almost the same with and without a nucleant. This suggests that when nucleation occurs it is quite rapid and that the time to observe a crystal is largely determined by the time taken for it to grow large enough to be visible. This time is not expected to be changed by a nucleant – which is what we observe. The growth rate of lysozyme crystals is known to decrease rapidly with decreasing supersaturation.25 Interestingly, Fermani et al.⁷ found that their nucleant (also based on gelatin) not only induced nucleation but greatly reduced the time taken to observe a crystal. We do not know why we find different behaviour, but we do note that they used a different protein, concanavalin A, and so it is possible that the difference may be due to the different proteins.

The TX-100 and particularly the gelatin buckypapers are effective nucleants, i.e., they induced crystallisation in metastable protein solutions. The ideal conditions for the growth of a well-ordered crystal are often at supersaturations that are too low to give rise to crystal nucleation. Such conditions are known as metastable, defined as those at which the drop will remain clear indefinitely if no nucleant, seed crystals or other nucleation enhancing procedure is applied. Thus our nucleants can be used to produce crystals at low supersaturations, allowing more ordered crystals to be formed.

Annealing the TX-100 buckypaper to remove the surfactant produced a buckypaper that is significantly more hydrophobic and has smaller pores; see Fig. 1. This buckypaper was ineffective as a nucleant; adding it to the crystallisation droplet did not reduce the minimum supersaturation at which crystallisation occurred. As both the surface areas and the pore sizes are changed by annealing, we cannot tell which one has the dominant effect in altering the lysozyme nucleation. It will require further systematic study to determine whether the ineffectiveness is the result of the change in hydrophilicity of the surfaces of the pores or of the change in mean pore size.
We found that a convenient form of buckypaper for crystallisation experiments is in small hair-like rectangular strips 0.2 mm by 1 mm. The data of Fig. 5A) were obtained with such strips. They were simply obtained using a sharp razor blade. With the gelatin buckypaper these strips often gave too many, too small, crystals all along their length. However, this problem was easily solved by simply using a smaller piece of buckypaper; see Fig. 5(B). Note that there the buckypaper strip is <100 μm by a little more than 100 μm, and that a large crystal has grown from it.

We also studied as nucleants both smooth surfaces coated with gelatin, and gelatin in solution, in order to compare their effectiveness as nucleants to that of gelatin-coated buckypaper. The trials were conducted at the lowest sodium chloride concentration that gave crystals with the gelatin buckypaper (3.6%). The gelatin-coated surfaces induced nucleation at low salt concentrations, as the gelatin buckypaper does, however we found them to be difficult to use. The gelatin-coated surfaces were hydrophilic, causing the droplets to spread out, which is undesirable. Also, a large and uncontrolled number of crystals appeared. We therefore abandoned studies of these systems. We also tried mixing a dilute (0.1 %) gelatin solution directly with the lysozyme and salt solution; see Fig. 5(C) for results. See the Materials and Methods section for details of these experiments. Here too, we found many small crystals at metastable conditions: the control droplets remained clear. The gelatin buckypaper is therefore more convenient to use and provides much greater control: it is more stable, i.e., does not have to be kept in a fridge, and by simply varying the size of the strip we could obtain the desired few, large crystals (compare Fig. 5(B) with Fig. 5(C), which shows the small crystals we obtained with the gelatin solution). Many small crystals are not useful for structure determination via X-ray diffraction so for this purpose our more controllable gelatin buckypaper is better than gelatin solution. However, gelatin solutions may still be useful for screening for conditions under which crystallisation occurs, and if a solid nucleant is undesirable for some reason, then gelatin solutions may then be very useful. Fermani et
al.\textsuperscript{7} have also studied gelatin-based nucleants for protein crystallisation. They studied the protein concanavalin A, and they found that their gelatin films were highly effective nucleants.

It may seem surprising that both our gelatin solutions and gelatin-coated surfaces induce the nucleation of lysozyme crystals, and that a third form of gelatin, a film, has also been found to be an effective nucleant\textsuperscript{7}. However, gelatin is largely collagen, and collagen exists as a triple helix. This has a diameter of 1.5 \textit{nm}\textsuperscript{20}, is quite rigid, and aggregates in solution (forming gels at concentrations higher than those we study). A recent study of recombinant collagen by Ramzi \textit{et al.}\textsuperscript{21} found that collagen helices existed as aggregates a few hundred nanometres across. We spun coated surfaces with very dilute gelatin solutions and studied the gelatin on the surface with AFM; see Fig. 4. On the surfaces were aggregates a few hundred nanometres across and the thickness of a single collagen triple helix high. In solution, an aggregate consisting of a network of collagen helices would essentially be a piece of collagen-walled porous medium a few hundred nanometres across. The gold nanoparticles aggregates studied by Hodzhaoglu \textit{et al.}\textsuperscript{11} may likewise resemble a porous medium. Thus our finding of nucleation due to gelatin alone, in solution as a film or on a smooth surface, is consistent with our hypothesis that materials with roughness on the length scale of the nucleus are good nucleants.

As the buckypaper strips are smaller than the desired crystals, observing crystallisation via optical microscopy is easy, despite the fact that the buckypaper is black. However, if required, large sheets of transparent buckypaper can be produced. See the Materials and Methods section for how this is prepared; we largely followed the work of Wu \textit{et al.}\textsuperscript{26} Figure 6 shows that lysozyme also nucleates on this transparent buckypaper, which is of course highly convenient for observing crystal growth via optical microscopy. On dried samples we can use SEM to search for crystals too small to be visible via optical microscopy. See Fig. 6(B) for an example. The density of such small, \textit{\sim} 100 \textit{nm}, lysozyme crystals on the surface of the buckypaper is relatively low, much less than one crystal per micron
squared. We do not know why the crystal did not grow larger; further study of this intriguing observation is left to future work.

C. Protein crystallisation trials: More difficult to crystallise proteins Lysozyme crystallises easily\textsuperscript{23}, therefore we also studied the crystallisation of proteins that are more difficult to crystallise, although here we were not able to obtain quantitative data on the crystallisation times. We worked with trypsin (from bovine pancreas, Sigma), the C1 domain of human cardiac myosin-binding protein-C (MyBP-C, supplied by Dr C. Redwood of Oxford University) and the non-structural protein 9 of the transmissible gastroenteritis virus (NSP9). Trypsin is a protease, i.e., a protein enzyme that breaks down other proteins, that is widely used in biotechnology. MyBP-C and NSP9 are both proteins relevant to human health, to cardiac disease and viral infection, respectively. The crystallisation conditions are described in the Materials and Methods section. MyBP-C and NSP9, due to their scarce supply, were not tested with the gelatin buckypaper. The MyBP-C was crystallised on an earlier TX-100-based buckypaper, which we used as pieces not strips.

Trypsin crystallises spontaneously at concentrations of 1.24M ammonium sulphate and above. At 1.20M ammonium sulphate, crystals grew on gelatin buckypaper within 3 days whereas controls as well as drops with TX-100 buckypaper remained clear for 9 days, after which crystals started appearing. At 1.16M ammonium sulphate, crystals only grew on the strips of gelatin buckypaper, all controls and TX-100 trials remaining clear. See Fig. 5(D) for a crystal of trypsin that has grown from our gelatin buckypaper.

MyBP-C is a much rarer and more difficult to crystallise protein. However, we were able to determine that 20\% PEG-3350 corresponded to a labile condition where all experiments including the controls resulted in clusters of rod-like crystals. 18\% PEG-3350 corresponded to metastable conditions. There, all controls remained clear whereas all experiments with buckypaper resulted in crystals with various
morphologies; see Fig. 7(A) for an example. Drops with 16% PEG-3350 remained clear. Before the development of the nucleant, hundreds of MyBP-C crystals were grown in clusters by conventional methods and X-rayed. The best resolution obtained from those crystals was 3.0 Å. By contrast, the best X-rayed crystals grown on the TX-100 buckypaper diffracted to a resolution of 1.6Å, almost twice as high. It is interesting to note that in one of these drops, containing 6 crystals, only one of the crystals was attached to the nucleant. That was the crystal that diffracted to the highest resolution. The other crystals in that drop diffracted with a resolution in the range 2 - 2.2 Å. Other drops (6 repeats) with nucleants present showed that in some drops the crystals were attached to the nucleant with no other crystals formed away from the it, and in other drops some crystals were attached to the nucleant and others were further away from it, but in all cases, the crystals in the drops containing nucleant were single, i.e., not in clusters. All the crystals belong to space group I4₁ with unit cell parameters a = b = 48.85 Å and c = 95.13 Å.

In the case of NSP9, both 19% and 20% PEG-3350 gave labile solutions, where crystals appeared in the controls as well as in the drops containing buckypaper. The buckypaper enhanced nucleation, producing showers of crystals at these conditions. The 17% and 18% PEG-3350 conditions were metastable, producing rod-shaped crystals in all the drops that contained buckypaper; see Fig. 7(B), whilst the controls all remained clear. Drops set with 16% PEG-3350 and below remained clear. In summary, our nucleants have been shown to be effective for a range of proteins and pHs, corresponding to crystallisation conditions at pH 4.5 (lysozyme) and 7.3-7.5 (MyBP-C and the NSP9). They worked with both salt and polymer precipitants.

4. CONCLUSIONS

To conclude, we have used materials with nanoscale porosity/roughness to control nucleation so as to obtain one or a few large crystals from solutions at low supersaturations – as required for structure
determination via X-ray crystallography. Growth at low supersaturations is expected to lead to more ordered structures which diffract to higher resolution, and indeed this is what was found for the MyBP-C crystallised with our carbon-nanotube-based nucleants. We believe that the carbon-nanotube based materials with nanoscale pores have great potential. The different effectiveness of our TX-100 and gelatin buckypapers implies that changing the surface chemistry and porosity changes the ability of the material to induce nucleation. Gelatin by itself was shown to promote nucleation, in accordance with a previous study\textsuperscript{7}. However, its use as a coating on buckypaper provided a much greater degree of control over the crystallisation process than when it was used either in solution or coating a flat film. Carbon nanotube films made with other dispersants may be even more effective nucleants. For example, films have already been prepared using dispersants as diverse as sodium dodecyl sulphate (SDS)\textsuperscript{14,15} and single-stranded DNA\textsuperscript{27}. These methods produce negative charges on the surfaces, which may be especially powerful for the crystallisation of positively charged proteins. Finally, nanotube films may also function as nucleants for systems other than protein solutions, for example in solutions of pharmaceuticals\textsuperscript{28}. Therefore, we believe that future work should test our materials as nucleants in other important systems where controllable crystallisation is desired.

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Figure 1. Characterisation of the buckypaper made with TX-100. (A) BET pore size distributions of buckypapers. (B) SEM (Hitachi S4000) image of unannealed buckypaper (t = 40 minutes probe sonication time). (C) and (D) are water droplets on buckypaper. (C) and (D) are for unannealed and annealed buckypaper, respectively, and the contact angles are 87.6° and 114.5°.
Figure 2. Plots of the nitrogen physisorption isotherms for A) TX-100 buckypaper, and B) gelatin buckypaper. \( V_{\text{ads}} \) is the volume in cm\(^3\) of nitrogen physisorbed per gram of the buckypaper as a function of the ratio of the pressure \( P_s \) to the pressure at saturation, \( P_0 \), which is atmospheric pressure as the experiment is conducted in contact with liquid nitrogen at atmospheric pressure. The upward pointing arrows indicate the adsorption isotherm while the downward arrows indicate the desorption isotherm. Note that in (B) although the apparent \( V_{\text{ads}} \) is actually higher for adsorption than for desorption, the difference is actually less than our estimate of our accuracy with which we can measure \( V_{\text{ads}} \) – approximately 2 cm\(^3\)/g. Thus in (B) if there is hysteresis, it is too small for us to measure.
Figure 3. Characterisation of the buckypaper made with gelatin. (A) AFM (NT-MDT) image of the surface of the buckypaper. (B) SEM (Hitachi S4000) image of the surface of the buckypaper. (C) is a water droplet on the buckypaper. The contact angle is 89.8°.
Figure 4. (A) AFM (NT-MDT) image of a surface that has been spin coated with a gelatin solution. Note that the heights of the features are approximately equal to the diameter of the collagen triple helix, 1.5 nm, and that their lateral extent is typically a few hundred nanometres. (B) Height profile along the pale green line in (A).
Figure 5. (A) Plot of the time at which a lysozyme crystal is first observed, as a function of sodium chloride concentration. Crystals are 5–10 μm across when they are large enough to be first observed. The lysozyme solution concentration was 20 mg/ml. The buffer was 0.1M sodium acetate at pH 4.5. Each point is the average of five crystallisation experiments, error bars are the standard deviations. (B) Lysozyme crystals on gelatin buckypaper, at 20 mg/ml in 3.6% NaCl. (C) Lysozyme crystals also at 20 mg/ml in 3.6% NaCl, with a droplet of 0.1% gelatin solution added. (D) Trypsin crystals on gelatin buckypaper. 30 mg/ml trypsin, 1.16 M ammonium sulphate, and 100 mM Tris pH 8.2.
**Figure 6.** (A) Optical microscopy image of lysozyme crystals on a sheet of transparent buckypaper. (B) SEM image of a lysozyme crystal too small to be visible via optical microscopy. The buckypaper is again transparent buckypaper.

**Figure 7.** Images of crystals growing from a TX-100 (t = 40) buckypaper nucleant at metastable conditions. (A) C1 domain of MyBP-C. (B) Non-structural protein 9 of the transmissible gastroenteritis virus (NSP9).

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