Effects of purification on the crystallization of lysozyme

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Abstract

We have additionally purified a commercial lysozyme preparation by cation exchange chromatography, followed by recrystallization. This material is 99.96% pure with respect to macromolecular impurities. At basic pH, the purified lysozyme gave only tetragonal crystals at 20°C. Protein used directly from the bottle, prepared by dialysis against distilled water, or which did not bind to the cation exchange column had considerably altered crystallization behavior. Lysozyme which did not bind to the cation exchange column was subsequently purified by size exclusion chromatography. This material gave predominately bundles of rod-shaped crystals with some small tetragonal crystals at lower pHs. The origin of the bundled rod habit was postulated to be a thermally dependent tetragonal \textsuperscript{\perp} orthorhombic change in the protein structure. This was subsequently ruled out on the basis of crystallization behavior and growth rate experiments. This suggests that heterogeneous forms of lysozyme may be responsible. These results demonstrate three classes of impurities: (1) small molecules, which may be removed by dialysis; (2) macromolecules, which are removable by chromatographic techniques; and (3) heterogeneous forms of the protein, which can be removed in this case by cation exchange chromatography. Of these, heterogeneous forms of the lysozyme apparently have the greatest affect on its crystallization behavior.

1. Introduction

The effects of impurities on the protein crystal growth process has attracted increasing attention [1–9]. It is now recognized that commercially available lysozyme is not pure, typically containing several macromolecular contaminants as well as assorted buffer and precipitant salts which together often comprise up to 15% of the dry weight [5,6,10]. It is also evident that simple recrystallization does not result in a markedly more pure protein; commercial lysozyme preparations are typically recrystallized from 3 to 6 times, yet still contain significant amounts of macromolecular impurities. Sources of the impurities may be from the added solution components or as a result of the preparation process. Alternatively, for many proteins, the growth affecting impurity may be an altered form of the protein itself. For example, proteases are often subject to auto degradation [8]. While the enzymatic function of lysozyme is the lysis of bacterial cell walls, its crystallization behavior can be affected by contaminating fungi [6].

Abergei et al. demonstrated that controlled contamination of turkey with other avian lysozymes affected the nucleation rate and crystal habit, with detectable cocrystallization [4]. Lorber et al. [5],
using electrophoretic and chromatographic methods, found impurities of \( M_r \approx 26, 45, \) and 80 K in various commercial lysozyme preparations. They also performed controlled contamination experiments and were able to correlate the amount of contaminating protein added to increased twinning. Wilson, using a differential dialysis system, was able to show the presence of five proteins in a commercial lysozyme preparation, having molecular weights of 14.6 (lysozyme), 17.5, 27.8, 46.8 and 77.9 K [10]. These molecular weights were maintained in the presence of disulfide reducing agents and a denaturing agent, indicating that all were single chain proteins and not aggregated lysozyme.

Impurities have been implicated in affecting the lysozyme crystal growth process. Lorber et al. indicated a correlation between the impurity level and the solubility, aggregation, and nucleation rate of tetragonal lysozyme [5]. Studies using Michelson interferometry suggested a significant impurity effect on the (101) face growth rate with the presence of a pronounced dead zone [7]. However, averaged face growth rate measurements, using chromatographically repurified protein, were found to be virtually identical to those obtained previously using material prepared by dialysis and recrystallization only [11,12].

Protein molecules are not rigid structures and another form of impurity could be protein which is in a different structural conformation. The evidence suggests that the thermally induced tetragonal \( \rightarrow \) orthorhombic transition for lysozyme is based upon a conformational change in the protein structure, and that the orthorhombic form is somewhat stable in the tetragonal phase region [13,14]. Structural shifts may also occur in proteins as a response to, for example, the solution pH, ionic strength, or binding to specific molecules in the solution. As in the case of the tetragonal \( \leftrightarrow \) orthorhombic shifts for lysozyme, these may be more rapid in one direction than the other, leading to mixed structural populations in apparently homogenous solutions. Thus, one must take the history of the protein into account when assessing purity. The most common method of purity assessment, gel electrophoresis of the reduced and denatured protein, relies on converting the protein into a random coil so that the electrophoretic mobility is only a function of the mass. In such a case, subtle structural changes which may be significant in terms of bioactivity or crystal growth are lost.

For the past several years, this laboratory has used only lysozyme which we have additionally purified by ion exchange chromatography for our growth rate experiments [11,12,15]. The initial impetus for the following work came from a reported 0.01 pH unit dependent change in the crystal habit for bovine pancreatic trypsin inhibitor [16]. A temperature, salt concentration, and pH dependent transition for lysozyme at acidic conditions has been shown on the basis of solubility studies [17]. We believed that a similar effect, with a tetragonal \( \rightarrow \) orthorhombic transition, might be found above pH = 7.0 based upon reports of the orthorhombic form occurring there when the protein is crystallized from egg white [14,18–22]. The appearance of well-formed tetragonal crystals instead, using highly purified lysozyme, throughout the range investigated (pH = 7.0–8.9) prompted the following study of the impurities in a commercial lysozyme preparation and the effects of their removal on crystallization behavior.

2. Materials and methods

Chicken egg white lysozyme from Sigma (St. Louis, MO, USA, grades I and III) was used throughout this work. Lysozyme concentrations were determined by UV absorbance, using an \( A_{281.5 \text{ nm}}^\text{lg} = 26.4 \) [23]. Measurements of lysozyme catalytic activity were performed using the \textit{micrococcus lysodeikticus} lysis assay [24].

The lysozyme was repurified by cation exchange chromatography as previously described [11], except that CM-Sepharose 6B-CL (Pharmacia) was used as the exchanger. Briefly, the protein was dissolved in and dialyzed against 0.1M sodium phosphate, 0.5% (w/v) sodium chloride, pH = 6.4, at room temperature. The dialyzed protein was clarified by centrifugation, then loaded onto a CM-Sepharose 6B-CL column (2.5 x 21 cm\(^2\)) equilibrated with the above buffer. The column was washed with equilibrating buffer and then eluted stepwise with 0.1M sodium phosphate, 2.87% sodium chloride, pH = 6.4. The column flow rate of \( \sim 4 \text{ ml/min} \) was maintained by a peristaltic pump. The eluted protein solution was brought to 10% sodium chloride by the slow addi-
tion, with stirring, of the solid salt, then cooled to 4°C to promote crystallization. The crystalline suspension was stored at 4°C until use, at which time the crystals were recovered by centrifugation, then dissolved in and dialyzed against the buffer to be used in subsequent experiments.

Size exclusion chromatography was performed with a Pharmacia FPLC system and a Superose 12 column at room temperature. The column was equilibrated in 0.2M ammonium carbonate buffer. The flow rate was 0.5 ml/min and fractions of 0.2 ml were collected. Protein was prepared for chromatography by dilution into the column buffer to a final concentration of ~1–5 mg/ml. The identity of the ovotransferrin and ovalbumin peaks was verified by running pure samples of both proteins (Sigma) through the column using the same conditions employed for lysozyme. Molecular weights of the impurities were further verified by SDS gel electrophoresis using a PhastSystem gel electrophoresis apparatus (Pharmacia), operated in accordance with the manufacturers instructions and using their low molecular weight standards calibration kit.

Crystallization experiments were set up using a Robotic Protein Crystallization System II (ICN Biomedicals, Inc., Costa Mesa, CA, USA) and CrysChem sitting drop crystallization plates (Charles Supper Co., Natick, MA, USA). Unless otherwise indicated, the final protein concentrations were 30 mg/ml, and the equilibration conditions for all crystallization solutions were 0.1M tris buffer (Sigma), 5% (w/v) NaCl (Sigma, analytical grade), with the pH ranging from 7.0 to 8.9 in 0.1 pH unit increments. The plates were sealed with clear tape and incubated at 20 ± 0.5°C. Crystal face growth rate measurements were performed using the techniques and instrumentation previously described [25,26].

X-ray crystallographic analysis of lysozyme crystals was performed using a Siemens multiwire area detector mounted on a Rigaku rotating anode X-ray generator. Cu Kα radiation was generated at 40 kV and 100 mA, and collimated with a graphite monochromator to 0.3 mm. Data was collected at a crystal to detector distance of 12 cm, with the detector −22°. The oscillation angle during data collection was 0.25° and each frame of data collected over 60 s. Data was reduced to structure factors with the Xegen package [27].

3. Results

The purity of the protein prepared by cation exchange chromatography was assessed with size exclusion chromatography. Previous electrophoretic analysis had indicated that the material was pure, with no other macromolecular contaminants detectable [11]. Size exclusion chromatography was employed to better quantitate any macromolecular impurities remaining. The column elution profiles at two detector scale ranges are shown in Fig. 1. Identical amounts of protein were injected onto the column in both runs. In the first profile, with the detector set at 2.0 absorbance units full scale, the purified lysozyme solution gave a single, fairly symmetrical peak. The slight tailing observed was due to lysozyme binding to the column, a common problem for lysozyme chromatography on carbohydrate matrix-based column media. This problem could be reduced somewhat by the use of higher ionic strength buffers. However, a balance must be observed between the buffer concentration and the protein concentration. Too high a buffer concentration would result in protein crystallization on the column bed. Initial attempts at sample preparation by the dialysis of concentrated protein solution into the 0.2M ammo-
Nium carbonate buffer resulted in showers of needle-shaped crystals.

A second profile, with the identical amount of protein loaded onto the column, but the detector set to 0.02 absorbance units full scale, displayed an almost flat base line until the onset of the lysozyme peak. The slight double peak ($\leq 0.0004$ absorbance units) immediately before the lysozyme peak was residual contaminating proteins. These contaminants have previously been identified as ovotransferrin and ovalbumin ($M_r = 80$ and 45 K) [5]. We further verified the correspondence of the contaminant peaks with these proteins by separate control runs with pure solutions of each protein. The ratio of the areas of the purified lysozyme peak at the 2.0 absorbance scale and the contaminants peak at the 0.02 absorbance scale was estimated by cutting out and weighing each peak. The slight double peak was divided at the lowest point between the maxima, and the corresponding areas of each part adjusted for the absorbance ratios of ovotransferrin or ovalbumin. On the basis of the adjusted area ratios the chromatographically repurified lysozyme was estimated to be $\geq 99.93\%$ pure with respect to detectable macromolecular contaminants. Trace amounts of a third contaminant, the serine protease inhibitor ovomucoid ($M_r = 28$ K) [28] cannot be estimated as the peak would be overwhelmed by the closely eluting lysozyme peak. However, ovomucoid binds less tightly to cation exchange columns than ovotransferrin or ovalbumin [29] and therefore it is unlikely that its residual concentration in the purified lysozyme would be greater than that of those two proteins. These results do not rule out the presence of other contaminants which do not absorb at 280 nm. However, the presence of such contaminants, assuming they are protein, has not been observed on SDS gels [5,10,17].

Fig. 2 shows the size exclusion column elution profile from the protein fraction which did not adsorb to the cation exchange column. This material contained the contaminating proteins present in the original lysozyme. In this particular case, most of the lysozyme bound to the column which resulted in the amount of contaminating proteins being greater relative to the amount of non-binding lysozyme. The four peaks, from left to right, are identified on the basis of their molecular weights as ovotransferrin ($M_r = 80$ K), ovalbumin ($M_r = 45$ K), ovomucoid ($M_r = 28$ K), and lysozyme ($M_r = 14$ K). The slight double peak found on the 0.02 absorbance units scale of Fig. 1 corresponds to the ovotransferrin and ovalbumin peaks of Fig. 2.

A series of crystallization trials, at 0.2 pH unit increments, over the pH range 7.0–8.9 were performed on the cation exchange purified lysozyme, the commercial protein, and fractions derived from it. These results are summarized in Fig. 3. The initial observation, that for purified lysozyme only tetragonal crystals were obtained from pH = 7.0 through 8.9, are shown in row (1). Columns (a)-(d) are the crystals obtained at pH = 7.4, 7.8, 8.2, and 8.6, respectively. A progressive increase in the size and reduction in the number of the crystals was observed as the pH of the crystallization solution increased. The crystals at higher pH values, as shown in panel (1d), were flattened along the four-fold axis, characteristic of tetragonal lysozyme grown at high supersaturations [30]. Crystals grown at lower pHs were more elongated along the four-fold axis, indicative of growth at lower supersaturations. The same results shown in row (1) were also obtained when the crystallization buffers were 0.05M sodium phosphate.
pH = 5.7–8.0 and 0.1M sodium bicarbonate buffer pH = 9.2–9.8.

X-ray crystallographic analysis of the tetragonal crystals grown at pH = 8.2 indicated a space group of $P4_{3}2_{1}2$ with unit cell parameters of $a = b = 79.1$ and $c = 38.1$ Å. These results are in close agreement with those found by previous analysis of the tetragonal crystal at acidic pH [22]. A crystal of dimensions $0.5 \times 0.4 \times 0.4$ mm$^3$ was used, yielding 7374 unique (or approximately 90% of the total possible) reflections at 2.0 Å resolution. The average redundancy was 4.7, giving an $R_{sym} = 6.9\%$. A more detailed study is currently underway to identify what structural differences, if any, exist between the acidic and basic crystalline protein structures.

The second row of Fig. 3 shows the crystals obtained with commercial lysozyme prepared by simply dissolving the protein from the bottle into distilled water. A sharp change was found in the crystal habit, occurring between pH = 7.6 and 7.8, with the crystals going from the characteristic tetragonal form to a bundled rod habit. Because of their small size, we were not able to confirm their type by X-ray diffraction. Presumably they are the same as the orthorhombic form previously obtained at basic pH [14]. The third row of Fig. 3 shows the results obtained when the protein was extensively dialyzed against distilled water prior to crystallization. In this case, only bundled rods were found. As in row (2), the rods appear to become finer with increasing pH.

The procedure for the cation exchange purification of lysozyme used a column equilibration buffer...
of 0.1M sodium phosphate, 0.5% sodium chloride, pH = 6.4. Following loading of the protein onto the column it was briefly washed with equilibrating buffer prior to eluting the bound lysozyme. This procedure often resulted in a large portion of the protein, up to 50% as determined by UV absorbance, passing through the column during the loading and washing steps. Electrophoretic analysis had previously shown that some of the non-binding protein was lysozyme, also shown by the size exclusion column elution pattern in Fig. 2. When rechromatography of this material was tried it typically again passed directly through the column. Separate experiments to determine if the non-binding lysozyme was due to overloading of the column indicated that this was not the case. In practice, only ~1/5–1/4 of the total lysozyme binding capacity of the column was used for purification runs (data not shown). Purification of the non-binding lysozyme away from the high molecular weight components was accomplished by FPLC on Superose-12. Crystallization of this purified non-binding lysozyme at basic pH again resulted in crystals of the bundled rod habit as shown in Fig. 3, row (d).

Activity measurements performed with the run-through, compared to the purified, lysozyme solutions had shown that the catalytic activity/mass of

![Fig. 4. Effect of the storage temperature on the face growth rate of tetragonal crystals at 20°C.](image)

Legend: (○) protein stored at 4°C; (×) protein stored at 37°C.

![Fig. 5. Effect of storage temperature on the nucleation and growth of crystals at 20°C. A 30 mg/ml protein solution was divided in half, with one aliquot stored at 4 and the other at 37°C. Mixtures were then crystallized by the sitting drop technique. The bar in (F) is 1.0 mm.](image)

Legend: (A) 100%, 4°C; (B) 50/50, 4/37°C; (C) 30/70, 4/37°C; (D) 20/80, 4/37°C; (E) 10/90, 4/37°C; (F) 100%, 37°C.
protein remained unchanged (data not shown). It was suspected that the origins of the non-binding protein, and thus the altered crystallization behavior, was due to the presence of lysozyme molecules with thermally induced altered conformations. However, warming an aliquot of the protein to 37°C for 4 h, followed by fractionation on a small CM Sepharose column with a shallow salt gradient, failed to conclusively demonstrate that this had occurred or, if it had, that the two conformations could be chromatographically resolved (data not shown).

Additional tests were made on whether heat treatment of the purified protein could affect its crystallization behavior. Comparative face growth rate measurements were made at 4 and 20°C, using protein solutions maintained at 4 and 37°C. The results for the 20°C growth rate measurements are shown in Fig. 4, and indicate that the storage temperature did not affect the growth rates. The results from the growth rate experiments at 4°C (data not shown) were comparable to the 20°C experiments.

To further test the effects of heat treatment on crystallization behavior, sitting drop experiments were set up to determine the effects on nucleation. As with the growth rate measurements, a lysozyme solution was divided into two aliquots, with one stored at 4 and the other at 37°C. Sitting drop crystallization experiments were set up at 0.1M sodium acetate pH = 4.6 and 0.1M tris HCl pH = 8.2, both at 5% NaCl and 20°C, using protein stored at the two temperatures and systematic mixtures of the two solutions. The results were similar at both pH values, and those for the pH = 8.2 experiments are shown in the photographs in Fig. 5. In most of the crystallization wells only tetragonal crystals were obtained, although a few crystals in the bundled rod habit were obtained at pH = 8.2. The data indicates that there was a pronounced effect of the storage temperature on the subsequent nucleation rate at 20°C. A systematic decrease in the number of nuclei formed was found with increasing percentages of the 37°C stored protein. However, only a slight increase in size is found with this decrease in nucleation rate, and the axial ratios are similar in all cases. This indicates that all crystals grew at approximately the same supersaturation. For comparison, note the pronounced differences with pH in the crystals of purified lysozyme in Fig. 3, row (1).

4. Discussion

The separation of crystal growth effects due to the presence of impurities versus those due to the solution behavior of the macromolecule itself have become important. A series of publications have recently shown that commercially prepared lysozyme has macromolecular impurities [5,6,10], that added contaminants affect the lysozyme crystallization process [4,5], and that lysozyme crystallization can be affected by microbial contamination [6]. We have also found macromolecular impurities to be present, which determination initially led to the use of cation exchange repurification of commercially available protein [11]. By using this procedure, it is possible to rapidly purify lysozyme in 3–5 g quantities to ≥ 99.9% purity with respect to other macromolecules.

Protein prepared by this procedure has been routinely employed in all growth rate experiments in this laboratory for the last several years [11,12,15]. However, previous to the work given above, we had not observed any appreciable difference in the crystallization behavior or growth rate data at acidic pH when compared to earlier experiments using commercial protein prepared only by recrystallization and dialysis. It was only when we searched for a pH dependent tetragonal → orthorhombic phase change above pH = 7.0 that we became aware of an effect which could be linked to the presence of impurities.

During the cation exchange purification process, a considerable fraction of the lysozyme loaded onto the column did not bind, but simply passed on through with the contaminating proteins. Control experiments indicated that the column had not been overloaded. Further, the percentage of non-binding material was not a function of the amount loaded onto the column, but was dependent upon the commercial preparation. Not all commercial lysozyme preparations are equivalent, as previously shown by others [5,10]. It has been this laboratories experience that there is variability even with the same material (different lot numbers) from a given vendor. Typically, with the lysozyme (Sigma cat. #L-7001) most commonly used in this laboratory ~ 10–50% does not bind to the cation exchange column. No apparent correlation exists between the amount of lysozyme which binds and the amount of other macromolecular impurities present. This suggests that the lysozyme
itself was affected by conditions during and after the original purification from the egg white. However, these changes in the crystallization behavior cannot be attributed to a tetragonal → orthorhombic conformational shift. Support for this comes from the crystallization behavior of the purified non-binding lysozyme and the purified binding lysozyme, and an inability to conclusively affect the crystal forms obtained at basic pH by heat treatment. A comparison of the lytic activity of the two protein populations also showed no significant differences. Thus, if there is a change in the protein it does not affect its biological function.

Cation exchange chromatography had been initially used to separate out non-binding lysozyme and other macromolecular impurities. Size exclusion chromatography was then used to separate the non-binding lysozyme from the other proteins. From a protein purification standpoint, the use of gel filtration is generally not a desirable method for the preparation of large quantities of material. This is especially true for lysozyme. Many of the matrix materials commonly employed for gel filtration resins are based upon carbohydrate backbones, which lysozyme binds. Agarose-based gel filtration media have been employed as an affinity material for the purification of lysozymes [31,32]. Thus, lysozyme tends to either elute at an abnormally low molecular weight (high retention volume) and/or have an elution profile which "tails". This tailing is evident in Fig. 1, despite the use of both high salt and pH buffer. However, this material had already passed through a cation exchange column without binding. Only a small amount of protein was required for the subsequent crystallization trials, therefore, we resorted to the use of gel filtration chromatography to achieve its purification. A high pH buffer was used to minimize protein interactions with the column matrix.

The use of cation exchange chromatography for lysozyme purification is not of itself novel. Early reports of lysozyme isolation from egg whites utilized adsorption onto bentonite [18]. The high isoelectric point of lysozyme suggested that weak cation exchangers would be an excellent method for its purification, and after their introduction they have been commonly employed to rapidly isolate lysozyme directly from egg whites [29,33–35]. Most likely, the ready availability of commercial protein preparations that have been purified by multiple recrystallizations has led to the neglect of this simple method of purification.

It is not surprising to find that macromolecular impurities persist even after multiple recrystallizations. The crystalline surface is covered with acidic and basic groups, and has a net charge which will be greater the further from the isoelectric point. Even at the isoelectric point there will be charged groups present. The putative macromolecular impurities also carry a net charge, and may bind to the crystalline surface which acts as an ion exchanger. Rapid nucleation and small terminal crystal size, typical of preparative recrystallizations, result in an increased surface area, all of which means an increased probability of binding other macromolecular impurities which may be present in the solution.

In experiments to test whether exposure to high temperatures (37°C) may be responsible for the appearance of the bundled rod habit crystals at basic pH, we found that the nucleation process is indeed affected, but that subsequent crystal growth is not. However, no conclusive evidence was obtained that exposure to high temperatures results in the appearance of the (presumably) orthorhombic form at basic pH. The nucleation results differ from those previously obtained by Berthou and Jollès [14]. They found that if the protein solution was preheated at 40°C, then cooled and crystallization initiated at 20°C, only orthorhombic crystals were obtained, with an apparent nucleation rate faster than for the corresponding tetragonal crystals from non-warmed solutions. We find instead that the tetragonal form persists despite the warming, and that the nucleation rate is apparently reduced. Berthou and Jollès did find a reduction in the nucleation rate of the tetragonal crystals when the pre-warmed materials were subsequently cooled to 4°C, similar to our findings at 20°C. These results suggest that a conformational change is involved. However, if such a change is important to the nucleation of a particular crystal form it is apparently not important for subsequent crystal growth.

These results demonstrate that one cannot rely on a manufacturer’s claim of purity by repetitive recrystallization of protein for use in crystal growth studies. Protein crystal growth may be affected by the
dialyzable (small molecule) and non-dialyzable (macromolecular) components also present in the bottle. However, even after removal of these components, the protein is itself subject to heterogeneity which may have an even greater affect on its crystallization behavior than either other macromolecular or small molecule components. In this study, very little effect is found on the lysozyme crystallization behavior at basic pH when the macromolecular and small molecule contaminants are removed. The predominant effect may come from heterogeneous forms of the protein itself. We are currently working on isolating these forms, and identifying the alterations responsible for the changes in crystallization behavior.

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