Locations of Bromide Ions in Tetragonal Lysozyme Crystals

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(Received 27 August 1997, accepted 16 February 1998)

Abstract

Anions have been shown to play a dominant role in the crystallization of chicken egg-white lysozyme from salt solutions. Previous studies employing X-ray crystallography have found one chloride ion binding site in the tetragonal crystal form of the protein and four nitrate ion binding sites in the monoclinic form. In this study the anion positions in the tetragonal form were determined from the difference Fourier map obtained from lysozyme crystals grown in bromide and chloride solutions. Five possible anion-binding sites were found in this manner. Some of these sites were in pockets containing basic residues while others were near neutral, but polar, residues. The sole chloride ion binding site found in previous studies was confirmed, while four further sites were found which corresponded to the four binding sites found for nitrate ions in monoclinic crystals. The study suggests that most of the anion-binding sites in lysozyme remain unchanged even when different anions and different crystal forms of lysozyme are employed.

1. Introduction

Most proteins bind with counterions in solution. When these proteins are crystallized, many of these ions may be expected to remain bound to the protein. The tetragonal form of chicken egg-white lysozyme (CEWL), is usually crystallized at a pH between 4 and 5, which is well below its isoelectric point of 11.4. The protein has positive net charge in this pH range (Tanford & Fevold, 1946; Steinrauf, 1959; King, 1959). Crystal solubilities showed a similar trend with thiocyanate ions causing the crystals to be least soluble and chloride ions causing them to be more soluble (Ries-Kautt & Ducruix, 1989; Pusey & Munson, 1991). Recently, the locations of bound nitrate ions in monoclinic lysozyme crystals have been reported (Rao & Sundaralingam, 1996). Knowing the number and locations of anions bound to the protein in the different crystal forms may lead to an understanding of the anion-induced polymorphism of lysozyme crystals, and help our understanding of the protein-crystallization process itself.

Earlier studies of tetragonal CEWL have located some of the binding sites. In their pioneering study of the structure of lysozyme in the tetragonal form, Phillips and co-workers found only one bound chloride ion (Blake et al., 1967). This was located between two molecules related by 4l symmetry. Binding studies have also been performed with phenolsulphothalein dyes in order to understand the enzymatic activity of lysozyme (Krishnamoorthy & Prabhananda, 1982a,b). A crystallographic study revealed two large pockets on the protein surface near the active-site cleft to be the binding sites for these dyes (Madhusudan & Vijayan, 1992). In other physico-chemical (i.e. non-crystallographic) binding studies the number of anions found includes those present in the solvent channels, making it impossible to determine those rigidly bound only to the crystalline matrix (Palmer et al., 1948; Carr, 1953; King, 1959). However, these studies do suggest that there may be more anion-binding sites in tetragonal lysozyme than the single site found by Phillips and co-workers. Very recently, in a high-resolution crystallographic study, the presence of an intermolecular anion-binding site was confirmed along with a sodium ion-binding site (Vaney et al., 1996).

In this study we investigated the binding sites of chloride and bromide ions in tetragonal lysozyme crystals employing ion-substitution crystallization experiments. The crystals were analyzed by X-ray crystallography, with the anions located from peaks in the electron-density difference map. These locations were also compared with the anion-binding sites of lysozyme located in earlier studies.
Table 1. Summary of least-squares refinement parameters

| Reflections from 6.0 to 1.8 Å with I > σ(I) | 11075 |
| Crystallographic R factor | 0.193 |
| Rmerge factor | 0.240 |
| Percent reflections used to compute Rmerge | 80.0 |
| Number of protein non-H atoms | 1004 |
| Number of ordered water molecules | 85 |
| Number of bromide ions | 4 |
| R.m.s. deviation of bond distances (Å) | 0.012 |
| R.m.s. deviation of bond angle (°) | 2.2 |
| R.m.s. deviation of dihedral angle (°) | 22.3 |
| R.m.s. deviation of improper angle (°) | 2.1 |

Crystallographic R factor = \( \frac{\sum |F_o| - |F_c|}{\sum |F_o|} \)

2. Materials and methods

2.1. Protein preparation and crystallization

CEWL purchased from Sigma was repurified as previously described (Ewing et al., 1996). Tetragonal crystals were grown in sitting drops at pH 4.6 in 0.1M sodium acetate buffer. Lysozyme chloride crystals were grown from solutions having a final concentration of 30 mg ml\(^{-1}\) protein and 5% NaCl, while the bromide crystals were grown from 85 mg ml\(^{-1}\) protein and 0.5M NaBr solutions. All crystals were grown at 293 K.

2.2. X-ray diffraction data collection and structure refinement

X-ray diffraction data were collected at room temperature, ~297 K, with a Rigaku R-AXIS II imaging-plane diffractometer mounted on a Rigaku RU200 X-ray generator and processed with DENZO and SCALPACK programs. Reflections for lysozyme bromide crystals were collected to 1.8 Å resolution [98.2% complete, I/σ(I) = 8.1, Rmerge = 5.3%]. Those for lysozyme chloride crystals were collected to 1.7 Å [97.5% complete, I/σ(I) = 6.6, Rmerge = 6.4%].

Structural refinements were performed for the lysozyme bromide data, while the lysozyme chloride data were only used to compute a difference Fourier map to identify bromide-ion positions. Refinements began with the coordinates of entry 6LYZ (Diamond, 1974), obtained from the Brookhaven Protein Data Bank. The space group P4\(_3\)2\(_1\)2 and unit-cell parameters \( a = b = 79.0\), \( c = 38.14\) Å were used. A molecular-replacement search confirmed the position of the lysozyme molecule. Initially, simulated annealing (SA) calculations were performed with the program X-PLOR (Brünger, 1991). These refinements used the reflection data in the 6.0–2.7 Å range, with a starting temperature of 300 K which was gradually lowered to 300 K.

Positional and temperature-factor refinements continued with the 6.0–2.7 Å reflections and included higher resolution data in each successive cycle. The difference Fourier map \( F_o(CI) - F_o(Br) \) was computed at 2.3 Å resolution to identify bromide-ion positions. The refinement continued for the lysozyme bromide model with the addition of ordered water molecules from the 2.3 Å resolution data, which were assigned when \( F_o - F_c \) peaks greater than 3σ were within hydrogen-bond distance (<3.5 Å) of the lysozyme atoms and other previously determined solvent atoms. The final refinements included the 6.0–1.8 Å reflections.†

2.3. Comparisons with other lysozyme binding sites

The various hen egg-white lysozyme structures in the Protein Data Bank were analyzed for any reported anion locations and were visualized using the program QUANTA (Molecular Simulations Inc.). In particular, the anion-binding sites in lysozyme reported in four recent crystallographic studies were considered (Madhusudan & Vijayan, 1992; Madhusudan et al., 1993; Madhusudan & Vijayan, 1999; Madhusudan et al., 1993).

† Atomic coordinates and structure factors have been deposited with the Protein Data Bank, Brookhaven National Laboratory (Reference: IAZF).
Fig. 2. Stereo images of the four bromide-ion locations identified by large peaks (>7σ) in the electron-density difference map: (a) the intermolecular Hal site, showing residues from two lysozyme molecules related by 41 symmetry (residues from the second molecule are shown by thin lines), (b) the Hal2 site, (c) the Hal3 site and (d) the Hal4 site. The crosses designate ordered solvent molecules in the vicinity.
3. Results

3.1. Final models from diffraction data

A summary of the final refinements is shown in Table 1. Because of the differences in the number of electrons between bromide and chloride (36 and 18 electrons), and assuming that bromide and chloride ions occupy the same positions in the crystal lattice, the bromide positions should be readily identifiable from the $F_o(\text{Br}) - F_o(\text{Cl})$ map. Four bromide ions were located which showed peak sizes of 7σ or greater in the difference Fourier map, along with a relatively weak fifth possible position. One bromide position (designated as Hal1) is at the molecular-crystal interface (Fig. 1) and its electron density is the highest of the four. Hal1 is closely associated with Tyr23 OH (at a distance of 3.0 Å) and Asn113 ND2 (at a distance of 3.2 Å) in the symmetry-related lysozyme (Fig. 2a). Two water molecules are also close to Hal1. Three other bromide ions (Hal2, Hal3 and Hal4) are located in different pockets of the lysozyme molecule in the crystal lattice. Hal2 interacts with the main-chain N atom of Gly26 and Ser24 OG (Fig. 2b). Hal3 is associated with the basic side chains of Arg14, His15 and the main-chain N atom of Ile88 (Fig. 2c). Hal4 is also closely associated with a cationic side chain Lys33 (the NZ atom at a distance of 3.5 Å) (Fig. 2d); it also interacts with a bridging water molecule which, in turn, interacts with the main-chain N atom of Arg5.

The four bromide ions described above are located in the larger of the two lysozyme domains. There is a fifth possible position in the smaller domain (Fig. 3). This site is near Asn74 ND2 and Asn65 ND2 (3.6 and 4.0 Å distant, respectively) and has a relatively small peak size in the difference Fourier map (~5σ). Although it is retained here as a halide ion (Hal5) for reasons discussed in the following sections, there is less certainty that this is a valid anion-binding site. The site is assigned as a water molecule in the refined structure.

3.2. Other anion-binding sites

The site Hal1 corresponds to the sole chloride ion binding site reported by Phillips and co-workers (Blake et al., 1967). The earlier study reported a slightly different location (near Arg114 of one molecule and Ser24 in the other) and the reason for this discrepancy is not clear. However, our location agrees with that from more recent high-resolution studies (Vaney et al., 1996).

Six nitrate ion binding sites were reported in monoclinic lysozyme crystals, three sites for each molecule in the dimer asymmetric unit (Rao & Sundaralingam, 1996). Only four of these sites are unique, as two sites are the same for both molecules in the dimer. Three of these sites correspond to the Hal2, Hal3 and Hal5 sites in tetragonal lysozyme. Two nitrate-binding sites were found from the crystallographic study of low-humidity (closely packed) monoclinic lysozyme crystals (Madhusudan et al., 1993), one of which corresponds to the Hal3 site. Finally, in the binding study with the dyes bromophenol blue and bromophenol red, one binding site was found for each dye (Madhusudan & Vijayan, 1992). The bromophenol red binding site corresponded to the Hal4 site.

4. Discussion

The five anions found in this study are clearly not the only ones present in tetragonal lysozyme crystals. Earlier investigations using physico-chemical methods found many more anions (Palmer et al., 1948; Carr, 1953; King, 1959). The remaining ions are either bound to the crystalline protein or free within the solvent channels. When a bound anion is in contact with the free water present in large channels, it is free to move in the vicinity of a central position (Frey, 1994). Thus, the occupancy of this central position or binding site is likely to be low and will be difficult to establish by a crystallographic study. This is confirmed by the locations of the five anions determined in this study which are all in small pockets in the crystal, often adjacent to bound water molecules (Fig 2). The absence of free water in these small pockets restricts the freedom of movement of the anions.
anion ensuring a high site occupancy (Madhusudan et al., 1993; Frey, 1994), and resulting in high electron densities at these sites.

The anions observed in this study have been designated as halide ions, assuming that both Cl\(^-\) and Br\(^-\) ions occupy these sites. However, the electron-density difference maps only confirm the existence of Br\(^-\) ions, and Cl\(^-\) ions may not bind at all these locations. The lower electron density of Cl\(^-\) and the low overall binding affinity for both Br\(^-\) and Cl\(^-\) evident by the high concentrations needed to precipitate lysozyme from solution, make the direct observation of bound Cl\(^-\) difficult and its presence can only be inferred from these studies. The presence of both halide ions is assured only at the Hal1 site, because of earlier observations of a Cl\(^-\) ion at this location. (Blake et al., 1967; Vaney et al., 1996). Two of the halide ions found by this study, Hal3 and Hal4, are associated with basic residues. Hal3 interacts with Arg14 and His15 (Fig. 2c), making this site a highly basic one and one likely to be occupied by anions in all crystal structures of this protein. Nitrate ions were also found at this location in the two monoclinic structures of lysozyme (Madhusudan et al., 1993; Rao & Sundaralingam, 1996).

Hal4 interacts with Lys33, with a bound water molecule associated with the main-chain N atom of Arg5 (Fig. 2d) and with the nearby Arg73 (not shown). This ion has the lowest electron density of the four, suggesting a low site occupancy. Such a low occupancy is most likely to be caused by the relatively large size of this pocket, and it is possible that it binds only the Br\(^-\) ion and not the smaller Cl\(^-\) ion. Significantly, this is the only halide ion binding site in the tetragonal form of lysozyme that does not seem to have an equivalent nitrate ion binding site in the monoclinic form.

Hal4 has been identified as the binding site of the dye bromophenol red, which is a much larger molecule (Madhusudan & Vijayan, 1992). As the dye-bound lysozyme is no longer active against its natural substrate, this site has been suggested as being essential to enzymatic activity by binding to the peptide component of bacterial cell walls. The results here corroborate the importance of this binding site to negatively charged and partially negatively charged groups. The position of Hal4 corresponds very closely with that of the partially negatively charged O atom in the sulfate group of bromophenol red (Madhusudan & Vijayan, 1992). Clearly this site can bind other partially or fully negatively charged groups, such as the O atoms in the peptide component of bacterial cell walls.

The Hal1 anion is the only one located at the interface between lysozyme molecules, where it is close to Arg114 and associates with partially positively charged atoms on both molecules and bound waters (Fig. 2a). Earlier studies had also indicated that this site was a point of close contact between lysozyme molecules in the tetragonal form (Blake et al., 1967; Imoto et al., 1972; Nadarajah & Pusey, 1996) and this accounts for Hal1 having the highest electron density of the five anions located in this study. In fact, the interaction between the molecules at this contact is central to the growth process of tetragonal lysozyme crystals. A chain of these contacts forms strongly bonded helices along the 4\(\bar{1}\) crystallographic axes (Nadarajah & Pusey, 1996). Growth of the crystals proceeds by the formation of aggregates corresponding to the helix in solution, followed by their addition to the crystal face by facetted-growth mechanisms (Nadarajah et al., 1997). Thus, the Hal1 anion plays a central role in the formation of tetragonal lysozyme crystals by facilitating the formation of these aggregates in solution (see Fig. 1).

The above discussion suggests that the Hal1 binding site may be unique to tetragonal lysozyme. The intermolecular contact corresponding to 4\(\bar{1}\) symmetry is absent in the monoclinic form and such an intermolecular binding site is not possible. However, one of the NO\(_3\) ion-binding sites in monoclinic lysozyme is almost the same as the Hal1 site, with respect to one molecule only (Rao & Sundaralingam, 1996). It is near the residues Asn113 and Arg114 (cf. Hal1 site with respect to the second molecule in Fig. 2a). Thus, in the same way as the Hal2, Hal3 and Hal5 sites, the Hal1 can also be considered to have an equivalent anion-binding site in the monoclinic form of lysozyme.

The location of the halide ion Hal2 is somewhat unexpected. Although it does associate with two polar groups, the main-chain N atom of Gly26 and with the partially charged H atom in the OH group of Ser24, there are no positively charged groups in its vicinity in the binding pocket (Fig. 2b). The importance of this binding site in lysozyme is also suggested by the fact that an NO\(_3\) ion binds here in the monoclinic form (Rao & Sundaralingam, 1996). This may seem surprising, but recent studies of the electrostatic fields of proteins in solution suggests that the contributions of polar groups to the field are significant and may even exceed those of charged groups (Warshel & Aqvist, 1991; Yang et al., 1993). The location of Hal2 near polar, but uncharged, groups confirms their importance to the overall electrostatic field of lysozyme and in particular to anion binding.

Given its low occupancy, some justification is needed for considering the final halide ion binding site Hal5 (Fig. 3) to be occupied by an anion rather than a water molecule. Two reasons can be given. Firstly, there is a sodium ion-binding site at a distance of 8.5 Å (Vaney et al., 1996), which, while rather distant, may increase the chance that an anion site is in the vicinity. Secondly, a nitrate ion was found on this site in the monoclinic form (Rao & Sundaralingam, 1996), indicating that this may be an anion-binding site in tetragonal lysozyme as well. Nevertheless, even if a halide ion is present at this site in the tetragonal form, it is likely to be only weakly bound to the protein.
Thus, in the regular monoclinic crystal form of CEWL, the four reported nitrate ion binding sites all have equivalent sites occupied by halide ions in the tetragonal form. Among the remaining binding sites for anions and dyes, the close-packed monoclinic form binds a nitrate ion at a different, intermolecular site (Madhusudan et al., 1993). The absence of halide ions at this site can readily be attributed to the different packing of tetragonal crystals. The binding site for bromophenol blue in tetragonal lysozyme is not at an intermolecular site, but at a large basic pocket near the enzyme cleft. This pocket contains the basic groups Arg21, Lys96 and Lys97 which do not form hydrogen bonds, are accessible to the free solvent and are presumably still protonated. Arg128 is also located in this pocket. As discussed earlier for bromophenol red, the binding of bromophenol blue also inhibits lysozyme activity with its natural substrate (Krishnamoorthy & Prabhananda, 1982a,b). However, unlike the binding site of bromophenol red which is the same site as Hal4, no halide ions were observed at this site.

The absence of anions on this site may seem surprising given the fact that bromophenol blue carries two negative charges and that its binding site is a highly basic one. Bromophenol red, on the other hand, is an uncharged but polar molecule with a less basic binding site (Madhusudan & Vijayan, 1992). The binding site of bromophenol blue is located in a large channel with free water molecules, while Hal4 is located in a smaller pocket, although as discussed earlier the occupancy of the Hal4 site is also low. This suggests that any anion(s) bound to the bromophenol blue binding site will only have a low occupancy, resulting in low electron densities. Even the much larger bromophenol blue molecule only has an occupancy of 0.30 on this site (Madhusudan & Vijayan, 1992).

5. Concluding remarks
This study shows the effectiveness of using ion-substitution experiments with X-ray crystallography to determine the ion-binding sites of crystalline CEWL. Four likely and one possible halide ion binding sites were found for the tetragonal crystal form. One of these sites corresponded to the intermolecular chloride ion binding site shown by earlier studies. Four of these sites also corresponded to four nitrate ion binding sites found in monoclinic crystals. The remaining binding site was found to correspond to that for the large bromophenol red molecule in the tetragonal form. These correspondences strongly suggest that the anion-binding sites for crystalline CEWL remain unchanged even when different ions and different crystal forms are employed.

We gratefully acknowledge the assistance of Dr. E. Snell, NAS/NRC resident research associate, NASA/MSFC, in the preparation of this manuscript.

References