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aFaculty of Advanced Life Science, Hokkaido University, Sapporo 0600810, Japan, bDepartment of Biotechnology, Graduate School of Engineering, Nagoya University, Nagoya 4648603, Japan, and cSynchrotron Radiation Research Center, Nagoya University, Nagoya 4648603, Japan. Correspondence e-mail: nobuhisa@nagoya-u.jp

Use of longer-wavelength X-rays has advantages for the detection of small anomalous signals from light atoms, such as sulfur, in protein molecules. However, the accuracy of the measured diffraction data decreases at longer wavelengths because of the greater X-ray absorption. The capillary-top mounting method (formerly the loopless mounting method) makes it possible to eliminate frozen solution around the protein crystal and reduces systematic errors in the evaluation of small anomalous differences. However, use of this method requires custom-made tools and a large amount of skill. Here, the development of a device that can freeze the protein crystal semi-automatically using the capillary-top mounting method is described. This device can pick up the protein crystal from the crystallization drop using a micro-manipulator, and further procedures, such as withdrawal of the solution around the crystal by suction and subsequent flash freezing of the protein crystal, are carried out automatically. This device makes it easy for structural biologists to use the capillary-top mounting method for sulfur single-wavelength anomalous diffraction phasing using longer-wavelength X-rays.

1. Introduction

X-ray crystallography is an efficient method for determination of the three-dimensional structures of biological macromolecules, from which important information can be extracted. Over the past decade, protein crystallography techniques have become highly sophisticated and a high degree of automation has been adopted. As a result, X-ray crystallography has become a common tool for biochemists with an interest in structural biology. Single-wavelength anomalous diffraction phasing using S atoms in native protein molecules, designated S-SAD, is becoming a widely applied technique in protein crystallography. Using this method, it is not necessary to prepare labelled crystals, such as heavy-atom derivatives or selenomethionine-substituted protein crystals. Despite the early suggestion of its availability (Wang, 1985), limited success was reported because of difficulties due to the small anomalous signal of S atoms (Hendrickson & Teeter, 1981; Dauter et al., 1999; Liu et al., 2000; Bond et al., 2001; Cianci et al., 2001; de Graaff et al., 2001; Gordon et al., 2001; Yang & Pflugrath, 2001; Brown et al., 2002; Lemke et al., 2002; Li et al., 2002; Micossi et al., 2002; Debreczeni, Bunkóczi, Girmann & Sheldrick, 2003; Debreczeni, Bunkóczi, Ma et al., 2003; Ramagopal et al., 2003; Lartigue et al., 2004; Sekar et al., 2004; Lakomek et al., 2009).

In the case of S-SAD phasing, use of a longer wavelength is advantageous for detecting the anomalous signal of S atoms. However, the accuracy of the measured diffraction intensity decreases at longer wavelengths because of the greater absorption and scattering effect. Data collection environments and processing methods specialized for using longer-wavelength X-rays have been discussed (Weiss, Sicker, Djunic-Carugo & Hilgenfeld, 2001; Weiss, Sicker & Hilgenfeld, 2001; Yang & Pflugrath, 2001; Cianci et al., 2008). The standard cryogenic crystal mounting technique (Teng, 1990; Garman & Schneider, 1997) has been improved with a new method that removes the buffer around the protein crystal before flash freezing of the crystals (Kitago et al., 2005). This capillary-top mounting method, previously referred to as the ‘loopless’ method, has been used to solve 14 novel structures including unpublished results (Watanabe et al., 2005; Hiyama et al., 2006; Kitamura et al., 2008). To date, however, this method has been carried out manually. The processes of harvesting and flash freezing the crystal are performed using both hands, and the mouth is also used for cryo-solution aspiration. Hence, this manual procedure requires specific skills. Automation of aspiration through a micropipette has been proposed for the routine use of this method (Lavault et al., 2006).

Some automated crystal harvesting devices driven by remote-controlled actuators have been developed (Hiraki et al., 2005; Viola et al., 2007). However, the crystal harvesting
operation is quite complicated and it is not easy to automate. Therefore, in our method, the crystal harvesting process is performed manually using a micro-manipulator, and the subsequent procedures have been automated. These developments make it possible to use the capillary-top mounting method routinely for longer-wavelength S-SAD phasing.

2. Description and operation

The prototype device (Figs. 1 and 2) was designed as a semi-automated device for crystal handling to allow the protein crystal to be harvested from the crystallization drop with manual movement of a micro-manipulator, and it is subsequently possible to remove the buffer and flash freeze the crystal automatically. The 7200CR Dual Head Epoxy Die Bonder (West-Bond, Anaheim, CA, USA) was chosen as the base component of the device. A magnet to hold the cryo-pin is placed at the head of the mechanically linked X-Y-Z micro-manipulator of the bonder, and a cryo-pin with a loop can be put there as the standard goniometer head for cryo protein crystallography (Fig. 2). There is a through-hole at the centre of the magnet where a suction line for removing the solution is connected. This makes it possible to use a custom-made cryo-pin for the capillary-top mounting method (Watanabe, 2006). In order to facilitate harvesting of the crystal, the tilt angle of the cryo-pin can be adjusted arbitrarily with a motor using an optical encoder attached at the manipulator arm. The docked or resting position of the bonder is used as the crystal freezing position where the conventional cryo-cooler nozzle is placed. The flow of the cryo-stream is gated by an air-driven mechanical shutter similar to those used at many synchrotron facilities (Giraud et al., 2009). It is also possible to use a standard cryo-pin, if it is not necessary to use the capillary-top method.

When the operator has successfully harvested a protein crystal from the crystallization drop, and moves the manipulator up from the drop to the freezing position, a series of events controlled by a sequential timer are triggered. A solenoid valve for the suction line opens and the solution around the harvested protein crystal in the loop is immediately aspirated through the line. The shutter then opens and the cryo-stream flash freezes the crystal, the surrounding solution of which has been removed. Typically, the solenoid valve for the suction line opens 100 ms after the manipulator backs to the freezing position as shown in Fig. 2(b), and the cryo-stream shutter opens a further 100 ms later. The intervals between sequenced events can be set by the user. Once the crystal is frozen, it is possible to recover the cryo-pin using a conventional cryo-tong, and it can be stored in a liquid nitrogen dewar. Short videos showing these procedures can be downloaded as supplementary data. This prototype is currently installed at Hokkaido University.

These videos are available from the IUCr electronic archives (Reference: HE5462). Services for accessing these files are described at the back of the journal.
The loop of the custom cryo-pin for the capillary-top mounting method was also improved. The conventional nylon loop glued to the tip of the glass capillary (Kitago et al., 2005) was replaced with a microlithography shaped polyimide film (Fig. 3). Polyimide film has already been used as a loop for MicroMounts (Thorne et al., 2003) and LithoLoops (Molecular Dimensions, Suffolk, England, and Protein Wave Corporation, Kyoto, Japan). Using polyimide film has some advantages over the nylon loop, including mechanical stability, negligible X-ray diffraction and absorption, and higher productivity. To locate the loop centre on the capillary axis, which is important to make the protein crystal sit properly on the tip of the capillary as shown in Fig. 4, we use an S-shaped design. To make the loop structure with the S-shaped film, one side of the scaffold is attached to the capillary, and then the other side is glued to the opposite side of the capillary, as shown in Fig. 5.

**Figure 3**
The microlithography polyimide film with S-shaped design. (a) Photograph of the whole sheet. (b) Magnified view of the sheet indicated by the red circle in (a). The red rectangle corresponds to one loop. (c) Example of dimensions of one S-shaped film for a 1 mm-diameter loop.

**Figure 4**
Examples of a crystal frozen using the newly designed loop. A thaumatin crystal was used. One division of the scale is 50 μm.

**Figure 5**
Procedures for making the loop structure using the S-shaped film. (a) One side of the film scaffold is glued to the capillary, and then the other side is glued to the opposite side of the capillary, as shown in (b). The tip of the capillary protrudes slightly into the S shape, as indicated by the blue arrow in the photograph.
3. Test experiment

To demonstrate this system, a longer-wavelength X-ray experiment was carried out using a thaumatin crystal as a sample. Thaumatin from *Thaumatococcus daniellii* (T-7638; Sigma, St Louis, MO, USA) was dissolved in distilled water (30 mg ml\(^{-1}\)) without further purification. Crystals were obtained under the conditions described by Ko *et al.* (1994), using 0.1 M N-(2-acetamido) iminodiacetic acid (ADA) buffer pH 6.5 and 0.6 M sodium potassium tartrate. The reservoir solution had a volume of 1 ml, and the crystallization drop consisted of 2 μl of protein solution and 2 μl of reservoir solution. For the diffraction experiment, a crystal of about 0.2 × 0.2 × 0.4 mm was used. The crystal was quick-dipped in cryoprotectant solution containing 30% (v/v) glycerol before harvesting. It was then mounted by the capillary-top mounting method using the device and the custom cryo-pin. Diffraction data were collected up to a resolution of 2.17 Å using the in-house chromium X-ray system described previously (Kitago *et al.*, 2005; Watanabe *et al.*, 2005). A total of 720 images with 0.5° oscillation were collected with a crystal-to-detector distance of 80 mm. The collected intensities were indexed, integrated, corrected for absorption and scaled using *HKL-2000* (Otwinowski & Minor, 1997). The detailed diffraction data statistics are presented in Table 1. The positions of anomalous scatterers were located using *SHELXD* (Sheldrick, 2008) at a resolution of 2.4 Å after analysing the substructure structure factors using *SHELXC* (Sheldrick, 2008). The initial phases were estimated using *SHELXE* (Sheldrick, 2008) and improved by density modification with a solvent content of 0.52. The protein model including 4–205 of 211 residues with side chains could be built automatically by *ARP/wARP* (Perrakis *et al.*, 1999). Part of an electron density map and an automatically constructed model are shown in Fig. 6.

Recently, Warkentin & Thorne (2009) have demonstrated that careful removal of external solution around the crystal using hydrophobic oil, such as Paraton-N, makes it possible to freeze the crystal as-grown without cryoprotectant. We have also performed experiments to show the usefulness of our method for mounting crystals without cryoprotectant. Thaumatin powder was dissolved in 0.1 M ADA buffer pH 6.5. A reservoir solution including 1.5 M sodium potassium tartrate was used for crystallization following the method of Warkentin & Thorne (2009). Crystals were frozen by three different procedures: using the standard loop with mother liquor, using the standard loop with Paraton-N and using the capillary-top mounting method. Each crystal was washed with the reservoir solution before freezing, in order to remove precipitate surrounding the crystal. Typical diffraction patterns are shown in Fig. 7. Frozen crystals mounted using our capillary-top mounting method, like those mounted using Paraton-N, do not give ice rings. Four (two for each) data sets with a total oscillation range of 10° were collected for these two mounting methods using different crystals. The conditions for the diffraction measurements were the same as for the above experiment. As shown in Table 2, the crystals frozen using the capillary-top mounting method are of almost the same quality as those obtained using Paraton-N.

### Table 1

Diffraction data statistics for a thaumatin crystal.

<table>
<thead>
<tr>
<th></th>
<th>Paragon 1</th>
<th>Paragon 2</th>
<th>Capillary-top 1</th>
<th>Capillary-top 2</th>
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<tr>
<td>Space group</td>
<td>P4(_2)(_2)(_2)</td>
<td>P4(_2)(_2)(_2)</td>
<td>P4(_2)(_2)(_2)</td>
<td>P4(_2)(_2)(_2)</td>
</tr>
<tr>
<td>Cell parameters (Å)</td>
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<td>a = b = 58.4, c = 150.5</td>
<td>a = b = 58.1, c = 150.6</td>
<td>a = b = 58.1, c = 150.2</td>
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<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Oscillation width (°)</td>
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<td>0.5</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Resolution limits (Å)</td>
<td>50.0–2.17 (2.21–2.17)</td>
<td>50.0–2.17 (2.21–2.17)</td>
<td>50.0–2.17 (2.21–2.17)</td>
<td>50.0–2.17 (2.21–2.17)</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>99.6 (93.0)</td>
<td>99.6 (93.0)</td>
<td>99.6 (93.0)</td>
<td>99.6 (93.0)</td>
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<tr>
<td>Multiplicity†</td>
<td>14.2 (9.2)</td>
<td>14.2 (9.2)</td>
<td>14.2 (9.2)</td>
<td>14.2 (9.2)</td>
</tr>
<tr>
<td>Wilson B factor</td>
<td>28.0</td>
<td>28.0</td>
<td>28.0</td>
<td>28.0</td>
</tr>
</tbody>
</table>

† Using ‘Scale Anomalous’ option of *HKL-2000*.

### Table 2

Diffraction data statistics (Å, °) for crystals frozen using Paraton-N and the capillary-top mounting method without cryoprotectant.

<table>
<thead>
<tr>
<th>Samples</th>
<th>Paragon 1</th>
<th>Paragon 2</th>
<th>Capillary-top 1</th>
<th>Capillary-top 2</th>
</tr>
</thead>
<tbody>
<tr>
<td>Oscillation range (°)</td>
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<td>10</td>
<td>10</td>
<td>10</td>
</tr>
<tr>
<td>Oscillation width (°)</td>
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<td>0.5</td>
<td>0.5</td>
<td>0.5</td>
</tr>
<tr>
<td>Resolution limits (Å)</td>
<td>50.0–2.22 (2.26–2.22)</td>
<td>50.0–2.22 (2.26–2.22)</td>
<td>50.0–2.22 (2.26–2.22)</td>
<td>50.0–2.22 (2.26–2.22)</td>
</tr>
<tr>
<td>Completeness (%)</td>
<td>99.6 (93.0)</td>
<td>99.6 (93.0)</td>
<td>99.6 (93.0)</td>
<td>99.6 (93.0)</td>
</tr>
<tr>
<td>Multiplicity†</td>
<td>14.2 (9.2)</td>
<td>14.2 (9.2)</td>
<td>14.2 (9.2)</td>
<td>14.2 (9.2)</td>
</tr>
<tr>
<td>Wilson B factor</td>
<td>28.0</td>
<td>28.0</td>
<td>28.0</td>
<td>28.0</td>
</tr>
</tbody>
</table>

† Using ‘Scale Anomalous’ option of *HKL-2000*.

**Figure 6**

An electron density map and an automatically constructed model of the thaumatin crystal. The green mesh shows the electron density map contoured at the 1.0σ level calculated using initial phases determined by longer-wavelength S-SAD with the capillary-top mounting method. The automatically built protein model constructed by the program *ARP/wARP* is also shown.
4. Discussion and conclusion

Our capillary-top mounting method is a powerful tool for longer-wavelength S-SAD phasing. The semi-automated device described here has convenient features for crystal mounting that will allow structural biologists to use the S-SAD method with non-labelled protein crystals. The structure of the fluorescent protein KillerRed (Carpentier et al., 2009; Pletnev et al., 2009), the crystal dimensions of which are about $250 \times 250 \times 200$ $\mu$m, was solved independently by S-SAD using this mounting device (PDB code 3a8s, unpublished results). In addition to the absorption effect, this method can reduce the background noise coming from the frozen cryo-solution. This feature is expected to be applicable to the measurement of micro crystals and ultra-high-resolution X-ray diffraction studies. The bore diameter of the tip of the current handmade capillary is about 40–200 $\mu$m. With these dimensions, there is none of the trouble in removing the external solution that could be caused by the viscosity of the mother liquor. However, the current device is only suitable for crystals larger than 40 $\mu$m. In order to enable use for very small crystals growing in a viscose solution, we are planning to develop a dual capillary device, the conceptual diagram of which is shown in Fig. 8. The idea for the device is functional separation of the capillaries: an inner capillary for crystal holding and an outer capillary for removal of the external solution.

This semi-automated device will also expand application of the capillary-top mounting method. The process of finding a suitable cryoprotectant for freezing a protein crystal is sometimes difficult and time consuming. We have shown that thaumatin crystals prepared using almost the same conditions as used by Warkentin & Thorne (2009) could be mounted as-grown using our capillary-top mounting method without cryoprotectant. Using hydrophobic oil sometimes causes damage to protein crystals. As our method removes the external solution around the crystal by aspiration, it is not
necessary to use hydrophobic oil to remove the solution. Qi & Jiang (2010) have tested 16 lysozyme crystals and 12 were successfully mounted without the use of cryoprotectant. Further experiments will be necessary to generalize the applicability. It was not possible to successfully freeze thau-matin crystals grown from 0.6 M sodium potassium tartrate. However, we think that this method is worth attempting for those crystals for which no suitable cryoprotectant can be found.

This capillary-top mounting method also facilitates the automated centring of cryo-cooled crystals. It is sometimes quite difficult to identify the frozen crystal in the lens-shaped frozen liquid. However, a crystal mounted by this method is located easily because the protein crystal will be frozen without surrounding liquid (Lavault et al., 2006). This feature will also be useful to spectroscopic investigations of protein crystals, such as optical monitoring of enzyme kinetics in the crystalline state (De la Mora-Rey & Wilmot, 2007).

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References

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