An arrayed-window microfluidic device for observation of mixed nanoparticles with an X-ray free-electron laser

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Abstract
A microfluidic device was developed for coherent diffraction imaging using an X-ray free-electron laser (XFEL-CDI). Liquid samples, which are separately packed in reservoirs of the device, are mixed in a flow channel after an XFEL irradiation as a trigger. An XFEL-CDI experiment was conducted and images of gold nanoparticles in solution were obtained with a resolution of several nanometers from single-shot coherent diffraction patterns. We were able to capture the images of various structures of mixed nanoparticles in the flow channel, clearly demonstrating the function of the microfluidic device. The direct observation of structural changes in liquids will contribute to development of chemistry and biology.

1 Introduction
X-ray free-electron lasers (XFELs) are characterized by high peak brightness, short pulse duration, and high spatial coherence [1]. While X-rays from conventional sources provide damages to samples, a femtosecond XFEL pulse enables us to obtain damage-free single-shot diffraction pattern before sample is destroyed, which is called the “diffraction-before-destruction” scheme [2]34. In particular, the high peak brightness and coherence of XFEL facilitates us to obtain coherent diffraction patterns to analyze even non-crystalline samples [5]. Single-shot coherent diffraction imaging with XFEL (XFEL-CDI) has been used for structural analyses of varieties of nanometer-sized samples [6] 7 8. XFEL-CDI is particularly suitable for structural analysis of hydrated samples with high radiation sensitivity. For example, wet nanomaterials [9] 10 and live bacteria [11] 12 have been observed with XFEL-CDI. These studies have revealed the natural internal structure of samples without freezing or drying. The method also has the potential to capture instantaneous states during structural changes in liquids, which contribute to new discoveries in biology, chemistry and materials science.

Here, we report a proof-of-principle work on the observation of the instantaneous states of nanometer-sized samples in liquids by combining the XFEL-CDI technique with a microfluidic device. Microfluidic devices, which are assemblies of microfluidic channels and reaction containers fabricated using microfabrication technology, have been used for applications in biotechnology and chemical engineering [13] 14. In this study, we developed a new device which has microfluidic channels with a number of irradiation windows to allow efficient measurement of coherent diffraction patterns of samples in solutions with XFEL pulses. The channel of the device has two reservoirs to hold liquids separately. The liquids in the reservoirs flow into the channel when one of the windows is broken by an XFEL shot. We observed
various aggregated conditions of spherical and rod-shaped gold nanoparticles mixed in the channel.

2 Method

2.1 Experiment system

The experiment was performed at BL2 of SPring-8 Angstrom Compact Free Electron Laser (SACLA) [15] [16]. As shown in Fig. 1, the device containing liquid samples was placed in an instrument called MAXIC-S (an upgraded version of Multiple Application X-ray Imaging Chamber [17]) (see Fig. 2). An XFEL single pulse with a photon energy of 4 keV and a duration of less than 10 femtoseconds was focused on an area of approximately 100 nm × 100 nm. The distance between the detector and the sample is 0.32 m. The multi-port charge-coupled device (MPCCD) detector [18] with eight sensor modules has a square-shaped imaging area with about 100 mm per side. The maximum scattering angle from a sample is 8.9°, which gives a theoretical resolution of 1 nm per pixel in the reconstructed image. The coherent diffraction pattern has a Fourier transform relationship with the difference of the electron densities between the sample and the surrounding solvent. The image of the sample in solution is reconstructed by phase retrieval calculations [19] [20].

The microfluidic device shown in Fig. 3 is a square chip (23 mm × 23 mm × 600 µm) with ten straight channels. Each channel has a dimension of 17 mm × 0.5 mm × 12 µm, two reservoirs that contain micro-liter liquid samples, and 45 irradiation windows made of a 200-nm-thick film of silicon nitride. The window is a parallelogram of about 130 µm × 200 µm. The interval of each window is 300 µm. The liquid sample is first injected to the reservoir and sealed with a polyimide tape. In this initial condition, the liquids stay in the reservoirs due to the surface tension of the liquids. The device is then mounted on the sample holder in the vacuum chamber of MAXIC-S.

An important feature of the microfluidic device is to allow observation of reactions of liquid samples immediately after they are mixed. Figure 4a shows a cross-sectional view of the microfluidic device with an XFEL beam path. With the first XFEL shot, a window at the end of the channel is broken, causing the liquid samples to flow into the channel, due to the rapid reduction of the pressure in the channel. Since each window is destroyed after single-shot irradiation, we need to change the windows for each irradiation. By irradiating the neighboring windows while shifting the device, several patterns from samples in solution can be obtained with the second and following shots. As shown in the Fig. 4b, c, the windows are irradiated from the window number 45 to 1. By arranging snapshots of samples according to the window number, it is possible to capture structural changes of the samples in situ along the reaction time.

The flow velocity in the channels is estimated based on the following equation:

\[
\nu = \frac{\Delta p}{\mu L} \left( \frac{1}{12} - \frac{16h}{\pi w} \tanh \left( \frac{\pi w}{2h} \right) \right),
\]

where \(\nu\) is flow velocity, \(\Delta p\) is pressure, \(\mu\) is fluid viscosity, \(L\) is length of the channel, \(h\) is height of the channel and \(w\) is width of the channel [21]. In this experiment, \(\Delta p\) is 1.0 × 10\(^5\) Pa and \(\mu\) is 9.4 × 10\(^{-4}\) kg/m \(\cdot\) s. \(L\), \(h\) and \(w\) are 17 mm,
12 µm and 0.50 mm, respectively. The estimated flow velocity is 75 mm/s. Considering, this velocity and the interval of the windows, the time resolution of the obtained snapshots is 4.0 ms.

2.2 Fabrication of microfluidic device

Figure 5 shows an exploded view of the microfluidic device. Two chips were fabricated by processing 4-inch silicon wafers with a thickness of 300 µm and a plane orientation of (110). Since wet etching process proceeds in the direction perpendicular to the (110) surface of a Si wafer masked by the resist pattern with parallelogram shapes, the interval of the windows can be set to 300 µm, which improves the time resolution of the snapshots according to the Eq. (1) [22] 23.

Each silicon wafer had a 200-nm-thick silicon nitride layer on its surface. After spin-coating with the resist ZEP520A (ZEON Corp.) to a thickness of 500 nm, the wafers were patterned by electron beam lithography (F5112 + VD01, Advantest). Reactive ion etching was subsequently used to remove the silicon nitride. In the area where the silicon was exposed, the irradiation windows and solution reservoirs were formed by wet etching with 30% KOH aq. The patterned wafers were cut into 23 mm square chips by laser dicing (DFL7340, Disco). Finally, two chips were bonded together to make a single device. The patterned adhesive photoresist TMMR SA390N (Tokyo Ohka Kogyo Corp.) was used as the bond between the two chips and the side walls of the flow channels. The estimation of the actual flow velocity is difficult because photoresist thickness uniformity after spin coating was insufficient. The value of the height in the size of the channel has a range from 12 to 24 µm. According to the Eq. (1), the estimated time resolution also has a range from 4.0 to 1.0 ms.

2.3 Liquid samples

In this study, gold nanoparticles (Cytodiagnostics Corp.) in two different shapes were used as samples in solution. One type was spherical gold nanoparticles and the other was rod-shaped ones. The gold nanospheres had a diameter of 50 nm. The length and the diameter of a gold nanorod were 40 nm and 15 nm, respectively. The nanospheres, which were covered with carboxyl groups, had a negative charge on their surface. The nanorod particles were covered with amino-PEG, having a positive charge on their surface. Due to the opposite charges, the two types of particles formed aggregates when the two solutions were mixed. Figures 6a, b show scanning electron microscope (SEM) images of a gold nanosphere and gold nanorods, respectively. Figure 6c shows an SEM image of an aggregate of gold nanoparticles.

Drops of the liquid samples were injected in the liquid reservoirs of both sides of the device with a pipette just before the XFEL-CDI experiment. The reservoirs were then sealed with polyimide tape, as shown in the right side of Fig. 3. UV-curing resin was applied to cover the edges of the tapes and to fill any small gaps.

3 Results and discussions

Figures 7a, b show optical microscope images of the liquid reservoir of the device in the vacuum chamber, observed before and after XFEL irradiation. After the irradiation, the liquid sample was dissipated, indicating that the solution was triggered to flow into the channel by the XFEL pulse. Various types of coherent diffraction patterns were obtained by irradiating the samples in the channel. Figures 8a, b, c
show typical examples of the patterns. We used the relaxed averaged alternating reflections algorithm and the shrink-warp algorithm [20] 24 to retrieve sample images from the coherent diffraction patterns. We performed 50 runs of image reconstructions from different initial random seeds for each diffraction pattern. Figures 8d, e, f show the averages of the 50 reconstruction results from the diffraction patterns shown in Fig. 8a, b, c, respectively. The images of the particles were retrieved with half-period spatial resolutions of 4.0 nm, 6.1 nm, and 12.7 nm, estimated by the phase retrieval transfer function [25]. This resolution allowed us to distinguish between gold nanosphere and nanorod particles. The values of phase shift contrast of 100-nm-thickness layers of Au and solvent are $3.7 \times 10^{-1}$ and $3.0 \times 10^{-2}$, respectively. Thus, the phase shift contrast loss of Au nanoparticles by solvent is almost negligible. The coherent diffraction pattern of a nanosphere particle was concentric, as shown in Fig. 8a. An intense streak in one direction appeared in the coherent diffraction pattern of a nanorod particle, as shown in Fig. 8b. We obtained several patterns caused by

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Fig. 6 SEM images of (a) a gold nanosphere particle, (b) gold nanorod particles, (c) an aggregate of gold nanorod particles and a gold nanosphere particle

Fig. 7 Microscopic images of liquid reservoir (a) before and (b) after XFEL irradiation

Fig. 8 Coherent diffraction patterns of (a) single gold nanosphere, (b) single gold nanorod, and (c) an aggregate of gold nanoparticles. d, e and f are the reconstructed images from (a), (b), and (c), respectively
nanorods, in which each intense streak was oriented in a random direction. One of the patterns is shown in Fig. 8b. We also obtained complex diffraction patterns that are different from these two types of patterns, as shown in Fig. 8c. In the image retrieved from Fig. 8c, an aggregate of many nanorod particles was observed. The image displays not only the three bright rod-shaped particles but also the dim-contrast objects at the top, left, and bottom of the aggregate. We estimate that the center of the image looks brighter than the edges because of the Gaussian distribution of the illumination beam intensity. This image indicates that there was a nanosphere particle having a negative charge in the center of the aggregate. The nanorod particles were attached to the entire surface of the nanosphere particle. Aggregates of a nanosphere and nanorod particles were also observed with SEM of Fig. 6c. In both images of the aggregates shown in Figs. 8f and 6c, multiple nanorod particles are attached to a nanosphere particle. In Fig. 8f, however, nanorod particles uniformly covers the surface of a nanosphere particle in contrast to the aggregate of nanorod particles lying on the membrane surface in Fig. 6c. We estimated that the electrostatic aggregate was broken due to the weak bonding of nanoparticles and the effect of drying process, such as the surface tension of the liquids and the concentration change. This result indicates that the method using the microfluidic device allows us to observe samples in solution without the effects of drying.

In this work, the device was designed to observe time-series structural changes of nanometer-sized samples in solution. Although we obtained several images of particles in various aggregate forms, time-lapse observation of the aggregation process has not been achieved with these images. This is presumably due to the uncontrolled flow velocity and the insufficient amount of data. We need to optimize the concentration of the samples and the design of the device, for example, the cross-sectional area of the flow channel and the volume of the reservoirs.

4 Conclusion

We have developed a microfluidic device for XFEL-CDI experiments to observe nanometer-sized samples in solution. The device has microfluidic channels with two pairs of reservoirs and dozens of windows for irradiation with XFEL pulses for single-shot image acquisition. The first-pulse irradiation induces liquid samples held in the reservoirs to flow into the channels and start mixing. The experiments demonstrated the microfluidic device functioning properly, and several coherent diffraction patterns were acquired. Gold nanosphere particles, nanorod particles and aggregates of both were observed by phase retrieval calculations using the patterns. In the aggregates, nanorod particles were uniformly attached to the surface of a nanosphere particle. This state is different from that captured in SEM images. This indicates that the microfluidic device allows us to observe samples in their natural state. Our microfluidic device has the potential to enable time-lapse observations of the structural change process of nano-sized samples in liquid. Such observations will make an important contribution to the fields of chemistry and biology in the future.

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