Fabrication of X-ray compatible microfluidic platforms for protein crystallization

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A B S T R A C T

This paper reports a method for fabricating multilayer microfluidic protein crystallization platforms using different materials to achieve X-ray transparency and compatibility with crystallization reagents. To validate this approach, three soluble proteins, lysozyme, thaumatin, and ribonuclease A were crystallized on-chip, followed by on-chip diffraction data collection. We also report a chip with an array of wells for screening different conditions that consume a minimal amount of protein solution as compared to traditional screening methods. A large number of high quality isomorphous protein crystals can be grown in the wells, after which slices of X-ray data can be collected from many crystals still residing within the wells. Complete protein structures can be obtained by merging these slices of data followed by further processing with crystallography software. This approach of using an X-ray transparent chip for screening, crystal growth, and X-ray data collection enables room temperature data collection from many crystals mounted in parallel, which thus eliminates crystal handling and minimizes radiation damage to the crystals.

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1. Introduction

Microfluidic platforms have gained widespread use in diverse fields like chemical synthesis, enzymatic and DNA analysis [1,2], proteomics [3,4], point-of-care, medical and clinical diagnostics [5–7], energy conversion [8,9], and protein/pharmaceutical crystallization [10–14]. Microfluidics offers several advantages in terms of reduced sample size and easy preparation, fine control over transport phenomena on the microscale, ease of scalability, detection and sample analysis on a single, integrated platform [15,16]. A large fraction of microfluidic devices reported in the literature to date have been fabricated using poly(dimethylsiloxane) (PDMS) and glass, which allow for rapid prototyping [13,17,18]. Issues still exist, however, with respect to compatibility with different chemistries, temperature, and pressure variation, as well as amenability to integration of analysis techniques. Many alternative materials and combination of materials have been explored for the fabrication of microfluidic platforms with the goal to overcome these issues for specific applications [19,20]. Typical challenges encountered in these efforts are bonding of different materials, interfacing with ancillary equipment, and the assembly of chips comprised of highly integrated, complex designs.

The focus of this work is on the application of microfluidic platforms in structural biology, specifically for protein crystallography. Protein structure determination involves handling of small sample volumes, fine control over transport properties during crystallization, and requires the collection and analysis of X-ray diffraction data from the crystals formed [21–23]. A microfluidic approach has the potential to meet these requirements. For example, we recently reported a microfluidic array chip for solid form screening of candidate pharmaceuticals [14], but this chip lacks X-ray transparency. The first step in protein crystallization involves screening a protein against a wide range of precipitants to identify suitable crystallization condition(s). Once one or more conditions have been identified, traditional methods require the crystals to be manually harvested, cryo-cooled and mounted in an X-ray beam for structure determination, all one crystal at a time. These challenging tasks, especially when crystals are small and fragile, often damage the crystal, affecting the quality of data collected. Several of the microfluidic platforms reported in the literature have tried to address these challenges, however most of them still require the crystal to be manually harvested before data collection [12,13,24,25]. X-ray transparent platforms have also been reported, but they have been limited to relatively simple, single-layer designs [11,19,26,27] that do not take advantage of the integrated fluid handling capabilities of multilayer microfluidics or require cutting out the section containing the crystal from the whole chip for further analysis [28].

Here we report on the fabrication of a microfluidic platform that allows for (i) screening up to 100 crystallization conditions while consuming minimal amount of protein solution, (ii) on-chip X-ray data collection from the protein crystals grown, while (iii) retaining
the fluid routing and manipulation capabilities of integrated, valve-based high throughput microfluidic systems. This is achieved by assembling a thin hybrid microfluidic chip comprised of layers of cyclic olefin copolymer (COC) and PDMS. We validate the utility of this chip and its use by crystallizing the soluble proteins lysozyme, thaumatin, and ribonuclease A on-chip, followed by X-ray diffraction data collection from the crystals grown while they still reside within the microfluidic platform mounted as a whole in the X-ray beam.

2. Materials and methods

2.1. Protein solutions

Hen egg white lysozyme (Sigma) was dissolved in 50 mM sodium acetate (Sigma-Aldrich) at pH 4.6 with 20% (w/v) glycerol (Fisher Scientific) at a concentration of ~100 mg/mL. Lysozyme concentrations were determined by UV absorbance measurements (Lambda 650 UV−VIS spectrophotometer, Perkin Elmer) at 280 nm using an extinction coefficient of 2.64 mL/(mg cm) [29]. For proof-of-concept crystallography experiments precipitant solutions of 1 M and 2 M NaCl (Aldrich) in 50 mM sodium acetate, pH 4.6 with 20% (w/v) glycerol were prepared. For screening experiments Hampton Crystal Screen chemicals were used directly (Hampton Research).

Thaumatin from Thaumatococcus danielli (Sigma) was dissolved in 100 mM NaH₂PO₄ (EMD Chemicals) at pH 6.5 at a concentration of 82 mg/mL. The protein concentration was determined by UV absorbance measurements at 280 nm using an extinction coefficient of 1.25 mL/(mg cm) [30]. A precipitant solution of 30% (w/v) Na/K tartrate (Malinckrodt) and 20% (w/v) glycerol in 100 mM NaH₂PO₄ pH 7.0 was used [31].

Ribonuclease A (R-5500, Sigma) from bovine pancreas was dissolved in 100 mM sodium acetate at pH 4.5 at a concentration of 229 mg/mL. The protein concentration was determined by UV absorbance measurements at 280 nm using an extinction coefficient of 0.70 mL/(mg cm) [32]. A precipitant solution of saturated NaCl in 100 mM sodium acetate at pH 4.5 was used [33].

2.2. Fabrication and operation of microfluidic platforms

Silicon wafers were patterned using negative photoresist SU8-25 and SU8-2050 (MicroChem). The microfluidic platforms consisted of different layers of polymers. The control layer was fabricated with COC (TOPAS Advanced Polymers Inc., 4 mil). Patternning of the COC was performed via hot embossing at 175 °C (Tₑ = 50 °C), using a master made out of a high temperature epoxy resin (Conapoxy FR 1080, 83:100 hardener:epoxy, by mass) using a hot press (Carver hot press, model 3851-0).

The fluid layer was fabricated out of PDMS (General Electric RTV 615, Part A/B). Inlets were drilled using a 750 μm drill bit (McMaster Carr). Bonding of the fluid and control layer (see SI for further details) was done using a 1% (v/v) solution of 3-glycidoxypropyltrimethoxysilane (GPTMS, Sigma Aldrich) and a 1% (v/v) solution of 3-aminopropyltrimethoxysilane (APTMMS, Sigma Aldrich). The completed assembly was placed on a flat COC substrate prior to setting up of crystallization trials. The use of vacuum for the operation of the devices eliminated the need to bond the fluid layer irreversibly to the COC substrate.

All solutions were introduced on-chip by pipetting 2–4 μL of protein/precipitant solution on the inlet ports, then pulling the fluid into the chip by actuation of the appropriate valves by applying vacuum through a manifold (Cole Parmer) attached to a vacuum pump (GAST; Model DOA-P704-AA). The vacuum was applied onto the chip through 24 gauge AWG, thin walled PTFE tubing (Cole Parmer) connected to a block of PDMS, which was positioned over the appropriate control line inlet. Once the chip was filled, all the inlets were covered with Crystal Clear Tape (Hampton HR4-511) to prevent evaporation and the trials were incubated. Even though some water is lost by absorption into the thin layer of PDMS, crystallization trials could easily be performed for a week, in most wells for two weeks, before solvent loss started to affect the crystallization outcomes, for example by precipitation of amorphous material.

2.3. Visualization of crystallization experiments and setup of traditional crystallization trials

Crystallization experiments were set up and visualized using either a stereomicroscope (Leica MZ12.5) equipped with a digital camera (Leica DFC295) or a computer controlled imaging system comprised of an optical microscope (Leica Z16 APO) equipped with an auto-zoom lens (Leica 10447176), a digital camera (Leica DFC280), and a motorized x-y stage (Semprexl KL66) controlled by Image Pro Plus (Media Cybernetics). Periodically birefringent images of the wells were taken.

Traditional microbatch-under-oil crystallization trials were set up combining 2 μL each of protein and precipitant solutions in a Greiner well plate (Hampton Research) at room temperature. Crystals were harvested using crystal mounts (Mitegen). Crystallization trials of lysozyme were performed in traditional well plates, large well devices, and array chips at 4 °C or at room temperature. Crystallization trials for thaumatin and ribonuclease A were performed at room temperature.

2.4. X-ray data collection

The microfluidic chips were mounted on a standard magnetic goniometer mount (Hampton Research) with an attached metal tube into which a slit was cut. A set-screw was used to secure samples. Diffraction data was collected either at room temperature or under cryogenic conditions. Cryo-cooling of samples was achieved by direct immersion into liquid nitrogen. Various sample-to-detector distances were used based on the quality of the crystal present. Typically data was collected using 1° steps with a 1 s exposure at an X-ray energy of 12.7 keV (λ = 0.979 Å) at Argonne National Lab (LS-CAT). Data from multiple crystals in the microfluidic devices was collected over a range of 10° (−5° to +5° from the normal) and an optimal subset of the frames was subsequently merged to obtain a complete dataset. Bench-top diffraction experiments were performed at the George L. Clark X-ray Facility at the University of Illinois using a General Area Diffraction Detector System (GADDS; Bruker) equipped with a four circle diffractometer and HiStar multiwire area detector. A rotating anode generator (Bruker M18XHF22) operating at 40 kV and 60 mA was used with a graphite monochromator supplying a Cu Kα radiation beam (λ = 1.54 Å or 8.048 keV). The sample to detector distance is ~18 cm. Data was collected in a coupled mode where 2θ=2θ such that multiple frames could be collected over a wider range of 2θ. Typically two such frames were collected, spanning the range of 2θ from 0° to 40°, up to a resolution of 2.3 Å.

Analysis of X-ray diffraction data collected at the synchrotron was performed using HKL2000 software for indexing, refinement, integration, and scaling (HKL Research) [34]. Diffraction data collected at the University of Illinois was analyzed using GADDS software (version 4.1.08, Bruker AXS) and Topas 3 (Bruker AXS). Subsequent processing of crystallography datasets was done using the CCP4 suite of programs [35]. Electron density maps were displayed using COOT [36]. Molecular replacement [37] for lysozyme was done using PDB structure 193L as the model [38].
3. Results and discussion

3.1. X-ray compatibility and materials characterization

In designing an X-ray compatible microfluidic device for use in protein crystallography three main considerations with respect to the interaction between materials and X-rays must be taken into account: (i) attenuation and (ii) scattering of X-rays passing through device materials, and (iii) the strength of diffraction resulting from a crystal. Attenuation originates from the absorption of photons by the material, thereby decreasing the intensity of both the incident X-ray beam and the resulting diffracted X-rays. Scattering is an elastic redirection of photons based on the internal structure of the material and can affect the signal-to-noise ratio. The strength of the diffraction signal from a crystal is related to not only the degree of order within the crystal, but also the packing density and the size of the crystal [39–43].

Fig. 1a shows the variation of the linear attenuation coefficient with the X-ray energy. In general, the linear attenuation coefficient is lower at higher X-ray energy. At relevant energies for X-ray data collection (here 12.4 keV), COC attenuates X-rays seven times less than PDMS and SiO₂. Fig. 1b shows the transmission factor for COC and PDMS as a function of film thickness. A thickness exceeding 3 mm is typical for conventional microfluidic devices, at which point the transmission factor is less than 10% (Fig. 1b). However for the hybrid COC–PDMS device as reported here, the transmission factor was more than 90% (Fig. 1a, dotted line shows data collection energy of 12.4 keV). An analysis of both PDMS and COC shows that these materials produce a characteristic scattering pattern, seen in the form of rings in magnified diffraction data. However this scattering occurs at relatively low resolution and does not impact the collection of high resolution diffraction data (see SI for more details).

3.2. Design and operation of microfluidic devices

The microfluidic device presented here enables the screening of protein solution against different precipitants, and also allows for structure determination of the protein once a suitable crystallization condition has been found. The main challenge here is to create a microfluidic chip that is still capable of fluid routing and compartmentalization to allow screening of many conditions, while also being X-ray transparent.

3.2.1. Array chip design

As discussed above, we chose to replace the traditionally thick layers of PDMS with thin films of COC because of its lower X-ray attenuation profile and beneficial material properties. However, COC is not suitable for the integration of valves into a microfluidic device because it does not have the same degree of flexibility as PDMS, an essential property for typical pneumatic valves [44]. To retain the ability to route and compartmentalize fluids on-chip, we use a thin layer of PDMS sandwiched between COC layers. This hybrid approach eliminates the bulk of PDMS usually found in traditional devices. Using a 20 μm PDMS layer sandwiched between two COC layers has only a minimal effect on the transmission of X-rays at energies relevant for X-ray data collection (Fig. 1a).

The hybrid microfluidic chips are comprised of a thin PDMS fluid layer sandwiched between and bonded to a COC control layer and flat COC substrate (Fig. 2a). The control layer is fabricated via hot embossing against an epoxy master. This epoxy master is fabricated by first creating a photoresist-on-silicon master using standard photolithography, followed by replication of this silicon master into a PDMS mold, which in turn is replicated in epoxy [20]. Sheets of COC (100 μm) were patterned with this epoxy master by hot embossing at 175 °C (Fig. 2a). The COC control layers have 25 μm deep and 50 μm wide negative relief patterns as the control lines. Fig. 2c shows the fidelity of pattern transfer into COC. Inlet holes for actuation of valves were then drilled in the COC control layer.

The fluid layer is fabricated in PDMS using standard soft lithography [44], with the COC control layer being bonded to the 70 μm thick fluid layer before being lifted from its photoresist master [45]. A perspective view of this assembly is shown in Fig. 2b. Inlet holes for the fluid layer were then drilled through this composite assembly. Lastly, this COC–PDMS assembly was placed on a 50 μm thick COC sheet to close the fluid channels. Both the 24-well array chip (Fig. 2) and the larger, single well chip for proof-of-principle testing (Fig. 3) were fabricated in this way. More details on the fabrication and assembly procedures can be found in Section 2 and Supplementary Information.

The array chips consist of a series of separate half-wells for protein and precipitant solutions arranged in columns. Each of these individual wells is a separate crystallization trial (Fig. 3a) and is isolated from the rest of the wells using a series of normally closed valves (Fig. 3a, green and blue valves) [46]. Since these valves are closed at rest, the chip can be transported easily after filling without disturbing the crystallization trials. Each half-well contains 50 nl of solution and the entire chip uses just 1.4 μL of protein solution for a 24-well design. These adjacent half wells are separated by a normally closed valve that allows mixing between the protein.
and precipitant solution through free interface diffusion. The time required to mix two solutions in adjacent wells can be estimated using Fick's law, \( t = \frac{x^2}{4D} \), where \( x \) is the combined length along which the two solutions are mixing and \( D \) is the diffusivity of the diffusing species (protein in this case, approximately \( \sim 10^{-6} \text{ cm}^2/\text{s} \)). For a mixing length of around 1 mm, the time needed to mix two solutions is approximately 20 min. The mixing time can be adjusted for other protein solutions with different diffusivities.

### 3.2.2. Array chip operation

The two half-wells are filled independently of each other using dedicated valve lines (blue and green valves for the aqueous protein and precipitant solutions respectively; see Fig. 3a) for each set of half-wells arranged in columns. These valves are closed at rest thus isolating the various chambers. The mixing valves (pink in Fig. 3a) between adjacent half-wells enable the protein and precipitant solution to mix by diffusion. The entire filling process is illustrated in Fig. 3. First 2 \( \mu \text{L} \) of protein solution is pipetted over the inlet of the protein line and vacuum is applied to the protein valve line (blue) which allows for dead-end filling of the protein solution into the series of protein half-wells (Fig. 3b). Once the protein solution has been introduced, the protein valve set is closed, isolating the solutions in the protein chambers. Next, 1 \( \mu \text{L} \) droplets of precipitant solution are pipetted over the inlets of each of the six precipitant. Vacuum is then applied via the precipitant valve lines (green) and the precipitant solutions flow into the appropriate half-wells (Fig. 3c). After closing the precipitant valve set, thus locking the precipitants in their respective chambers, the inlets for filling and actuation are sealed with Crystal Clear Tape, preventing evaporation of the filled solutions. Next, the mixing valves (pink) located between the protein and precipitant chambers are actuated, allowing sets of two adjacent solutions to mix by free interface diffusion (Fig. 3d). We allowed the solutions to mix for 20 min for all experiments performed here. After mixing, the mixing valves are closed, the inlets of these control lines are sealed with Crystal Clear Tape, and the chips are incubated at either 4 °C or room temperature for 8–48 h. Previously, we have used a similar microfluidic array chip for solid form screening of candidate pharmaceuticals [14], but this platform lacked X-ray transparency.

### 3.2.3. Single-well chip design, operation, and proof-of-concept experiments

Before completing the full design and fabrication of the 24-well array chip, we wished to confirm that indeed high quality diffraction data can be collected from crystals residing inside a microfluidic well comprised of COC layers as well as a thin PDMS
layer (Fig. 4a). These chips for validating the X-ray compatibility of the materials are made using a similar fabrication scheme described earlier (Section 3.2.1). They consist of a large rectangular 2.36 mm × 3.36 mm × 25 μm wells with a volume of 0.2 μL. Six posts were fabricated within the fluidic well chamber to provide support for the thin PDMS membrane. The COC control layer consisted of an identical sized chamber aligned directly above the fluid well (Fig. 4a). Pre-mixed solutions of protein and precipitant (see Section 2) were introduced into the chip by pipetting a drop over the inlet of the fluid layer and pulling it into the fluid layer well by applying vacuum to the inlet of the control layer (Fig. 4a). The chip was allowed to incubate for 48–72 h.

To test the effectiveness of the chip materials for on-chip X-ray data collection, we first grew crystals of model proteins lysozyme (Fig. 4c), ribonuclease A (Fig. 4d), and thaumatin (Fig. 4e) in single well chips. The whole chips were then mounted onto a goniometer mount (Fig. 4b, see Section 2) and taken to Argonne National Lab (LS-CAT). Fig. 4c–e shows on-chip diffraction data collected from these chips from the different model proteins. Even though a scatter ring from the materials is present, good quality, circular diffraction spots are still clearly visible, sufficient for structure determination.

3.3. Screening and on-chip protein structure determination

Identification of suitable crystallization conditions by screening protein solution against a wide variety of crystallization mixtures (cocktail of salts, buffers, and precipitants) is the first step in the process of protein structure determination. Fig. 5 shows a 24-well array chip that allows for the testing of six different precipitant solutions in quadruplicate. A series of array chips was used to test each of the individual conditions multiple times and the results were compared to those obtained using a traditional microbatch wellplate.

The microfluidic array chips were validated for crystallization screening experiments by testing solutions of lysozyme against the 50 condition Crystal Screen kit (Hampton Research) at room temperature. After one week crystals were observed in 32 out of the 50 conditions in the array chips (Table 1) compared to only 26 hits in

![Fig. 3. Schematic of the design and operation of a 24-well hybrid array chip. (a) Complete chip with the fluid layer shown in black, and the various components of the control layer are shown in respective color according to function. The window structures are present to decrease the total material present in the path of the X-ray beam. (b) Blue valves are actuated to fill in protein. (c) Green valves are actuated to fill in precipitant. (d) Pink valves between the protein and precipitant chambers are actuated to allow diffusion between the chambers over time. (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)](Image)
Fig. 4. (a) Design of simple large well microfluidic chip. Numbers in brackets represent the total height of each layer. The feature size in both the COC and PDMS layers is 25 μm. (b) Optical micrograph of an X-ray transparent large well device mounted on a beamline at LS-CAT, Argonne National Lab with the inset showing a close-up of the modified magnetic goniometer mount on which the device is mounted. Optical micrographs of microfluidic chips with crystals of (c) lysozyme (d) ribonuclease A and (e) thaumatin. On-chip diffraction data (collected at LS-CAT, Argonne National Lab) from each of the crystals at cryogenic conditions is displayed alongside as well. Scale bars correspond to 300 μm.

Fig. 5. Optical micrograph of crystals of lysozyme grown on a 24-well hybrid array chip, with a magnified view of one of the crystallization chambers.

the microbatch wellplate. X-ray compatibility of the chip allowed us to easily identify whether the hits were indeed protein crystals or just salt crystals using a local X-ray source (Bruker APEX). A comparison of the results obtained in our well plates versus the traditional microbatch method shows that 21 of the conditions produced crystals on both platforms while 5 conditions yielded crystals uniquely in the microbatch wellplates and 11 hits were observed uniquely in the microfluidic chips. The variability between these results can be explained by (i) differences in the method of mixing of protein and precipitant solutions; (ii) slow concentration of solutions in the microfluidic chips over time; and (iii) the stochastic nature and variability of crystallization trials [47–49]. Note that typical

<table>
<thead>
<tr>
<th>Condition number</th>
<th>On-chip(^a)/off-chip(^b)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
</tr>
<tr>
<td>1–10</td>
<td>-/−</td>
</tr>
</tbody>
</table>
| 11–20            | +/+| +/−| −/−| −/−| −/−| +/+/+ | +/+ | +/+| +/+| +/+
| 21–30            | +/−| +/−| −/−| −/−| −/−| +/+/+ | +/+ | +/+| +/+| +/+
| 31–40            | +/−| +/−| −/−| −/−| −/−| +/+/+ | +/+ | +/+| +/+| +/+
| 41–50            | +/−| +/−| −/−| −/−| −/−| +/+/+ | +/+ | +/+| +/+| +/+

\(^a\) Crystallization trials set up on a 24-well array chip.
\(^b\) Trials set up on a microbatch tray under oil.
\(^c\) “−” indicates absence.
\(^d\) “+” indicates presence of crystals.
microbatch wellplate experiments need 2 μL of protein solution per condition whereas a whole 24-well array chip can be filled using less than 2 μL of protein solution.

**Strategy for on-chip data collection** – Once crystals have grown on-chip, we want to collect X-ray data with the crystals still residing in the microfluidic wells. In tradition crystallography, crystals are manually looped out, cryo-cooled, mounted in a cryo-stream, and rotated in the X-ray beam for data collection. The dimensions and design of the array chips makes rotation of the chip for the collection of a complete dataset very difficult. The chip is also too large to fit within typical cryo-streams on crystallography beamlines, necessitating data collection at room temperature.

The inability to utilize cryo-cooling to mitigate the effects of radiation damage [50] led us to adopt an alternative strategy where we collect small wedges of data (10⁶) from a large number of crystals at room temperature and later merge these wedges to form a complete dataset. The use of small wedges of data limits the time of exposure, and thus the extent of radiation damage. These wedges can be merged to form a complete dataset, provided that non-isomorphism between crystals is minimal and the orientation of the grown crystals is mostly random. This method has been used previously to obtain structural information from tiny or fragile crystals or crystals which suffer from excessive radiation damage [51,52].

The advantage of applying this data collection strategy to crystals grown in a microfluidic chip is the ease by which a large number of crystals can be generated. Furthermore, the fine control of transport on the microfluidic scale allows for improved reproducibility both in crystal quality and isomorphism between crystals grown in different wells or even in different chips. Analysis of lysozyme crystals grown on-chip demonstrated that 98% of the crystals (47 out of 48) showed a coefficient of variation (standard deviation divided by the average) in the unit cell dimensions of ~0.1%.

**Analysis of on-chip collected X-ray data** – We collected and analyzed room temperature X-ray data of lysozyme crystals grown in 24-well chips using the same crystallization conditions as used for the proof-of-principle experiments. For comparison, we also collected X-ray data at cryogenic temperatures on a lysozyme crystal grown in a wellplate and mounted using a loop with Table 2 showing a comparison of the important crystallographic statistics between the two methods.

Analysis of the crystallographic parameters and statistical measures of quality for both data collection and structural refinement via molecular replacement show that the quality of the data was not significantly affected either by collection at room temperature or because of the merging of data from multiple crystals. The diffraction data was analyzed over the range of 50–1.55 Å (the diffraction limit for the on-chip data) to enable direct comparison between the other parameters. Other data collection parameters such as Rsym, completeness, redundancy, and I/σ and structural refinement parameters such as Rfree, are comparable for the two cases, despite differences in crystal size, resolution, and signal-to-noise resulting from the presence of device materials. Interestingly, due the absence of both physical handling and cryo-cooling, the mosaicity for the room temperature merged data is nearly an order of magnitude lower (i.e., better) than the mosaicity of the single-crystal cryogenic datasets. Fig. 6 compares the diffraction patterns and electron density maps obtained from the traditional, single-crystal cryogenic data and the on-chip, multi-crystal, room temperature data. The high resolution data obtained in each of these cases is evident from the electron density maps of, e.g., aromatic amino acid side chains with the rings clearly visible.

### Table 2

<table>
<thead>
<tr>
<th>Parameter</th>
<th>On-chip⁴</th>
<th>Traditional (cryogenic)³</th>
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</thead>
<tbody>
<tr>
<td>Unit cell dimensions</td>
<td>a = b = 79.69 Å</td>
<td>a = b = 78.81 Å, c = 37.02 Å</td>
</tr>
<tr>
<td>Space group</td>
<td>P4₁,2₁,2</td>
<td>P4₁,2₁,2</td>
</tr>
<tr>
<td>Observations (unique)</td>
<td>783,994</td>
<td>223,433 (17,510)</td>
</tr>
<tr>
<td>Resolution</td>
<td>50.1–1.55 Å</td>
<td>50–1.55 Å</td>
</tr>
<tr>
<td>Rsym</td>
<td>0.064 (0.362)</td>
<td>0.052 (0.102)</td>
</tr>
<tr>
<td>Mosacitxy</td>
<td>0.03–0.08</td>
<td>0.21–0.34</td>
</tr>
<tr>
<td>Redundancy</td>
<td>22.9 (5.7)</td>
<td>7.7 (7.7)</td>
</tr>
<tr>
<td>Completeness</td>
<td>98.1% (83.4%)</td>
<td>99.7% (100%)</td>
</tr>
<tr>
<td>l/σ</td>
<td>51.4 (3.9)</td>
<td>42.3 (19.4)</td>
</tr>
<tr>
<td># of frames</td>
<td>363</td>
<td>100</td>
</tr>
<tr>
<td>Refinement R(Rfree)⁵</td>
<td>0.164 (0.227)</td>
<td>0.173 (0.276)</td>
</tr>
</tbody>
</table>

**Notes:**

⁴ Merging of small datasets from “multiple crystals” analyzed on-chip within a 24-well device at room temperature.

⁵ The “traditional” sample was grown using microbatch techniques and mounted using a standard crystal mount for cryogenic data collection. Reported values are for all hkl’s. Values in parenthesis represent the value for the highest resolution shell except where indicated for the number of observations as compared to unique reflections and R(Rfree) for the Ramachandran statistics where the number in parenthesis indicates the number of residues in a given region. Data was analyzed over the range of 50–1.55 Å to enable a direct comparison with the data collected from merging the diffraction data taken from multiple crystals at RT.

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![Fig. 6](https://example.com Fig6.png)

*Fig. 6.* X-ray diffraction data collected at LS-CAT, APS from (a) a cryo-cooled lysozyme crystal grown using the traditional microbatch method and mounted on a standard crystal mount, (b) a lysozyme crystal on a hybrid array chip at room temperature. Corresponding figures below show an omit map for the Trp 111 residue, the calculated map is displayed in blue (σ = 2.0) and the difference map is in green (σ = 3.0). (For interpretation of the references to color in this figure legend, the reader is referred to the web version of the article.)
4. Conclusions

In summary, the microfluidic platform reported here allows for crystallization screening and subsequent on-chip X-ray data collection of crystallized proteins. The fabrication scheme uses a combination of COC and PDMS layers, and yields a platform that retains fluid routing capabilities via pneumatic valving while also achieving X-ray transparency. The dense network of integrated pneumatic valves allows for fast and easy setup of the crystallization screens, which in turn facilitates the reproducible growth of a large number of isomorphous crystals. The valves are only opened by vacuum actuation during filling and mixing. The valves are closed in rest, so the individual crystallization trials are isolated during crystallization and data collection in the absence of connections with ancillary equipment. The whole chip can be transported to and mounted in the X-ray beam for collection of diffraction data wedges from many crystals still residing on chip at room temperature. By merging these wedges of data a complete data set for full structure determination is obtained. This approach eliminates the need for crystal harvesting and cryo-cooling, and simultaneously avoids the detrimental effects of the radiation damage associated with room temperature data collection.

The screening capabilities of the platform were highlighted using a standard screen, followed by comparison of the results with the same screen performed via a traditional crystallization method. The platform was validated by collecting data from several proteins like lysozyme, thauaminit, and ribonuclease A. By solving the structure of lysozyme using only diffraction data from many crystals residing on-chip we confirmed that the strategy of merging wedges of data from many crystals to obtain a complete dataset can be applied to these microfluidic platforms.

Going forward, the methodology reported here can be used to solve structure of novel proteins, especially those that (i) yield small, fragile crystals, (ii) have no known cryogenic crystallization conditions, and/or (iii) have been known to yield low resolution data using traditional cryogenic data collection. In addition to allowing for the routine collection of static structural information of proteins at biologically relevant temperatures as reported above, the fine control over transport phenomena coupled with on-chip analysis capabilities offered by these microfluidic platforms could enable a range of other biological studies. For example, precise microfluidic fluid handling could be utilized to enable dynamic structural studies based on the addition of various stimuli such as ligands, electrochemical agents, acid/base to affect the pH, or even changes temperature. Such studies have the potential to shed light on structural changes associated with protein function in unprecedented ways.

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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.snb.2012.08.048.

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