

4. CRYSTALLIZATION

grill into a square Petri dish before pouring the agar, and the grill remains embedded in the media. Small glass beads can be added to each segment of the 48 grids to spread the solution. Colonies are picked by a colony picker, and are grown in 96-well deep-well blocks containing media and appropriate antibiotics. Using a liquid-handling robot and a colony picker for a single round of cloning, it is possible to achieve >80% efficiency in a 96-well format. The same robotic setup can be used for the subsequent recovery of recombinant plasmid before DNA sequencing and protein expression.

4.4.4. Protein expression and purification

Small-scale protein expression in an HT format enables evaluation of the expression, solubility and purification of target proteins or multiple constructs. Autoinduction allows many protein expression tests in parallel without having to monitor cell densities to optimize induction, and is thus ideal for HT approaches. Each protein in a 96-well plate will have different chemical properties and sizes, and thus affinity tags are the method of choice for parallel purification of multiple proteins on a single platform. Dot blots can be used to test expression or solubility, and miniaturized resin deposited in a 96-well plate can be used for a purification test. Although up to 90 µg of protein have been purified from 1 ml of *E. coli* cell culture (Scheich *et al.*, 2003), the amount of protein obtained from small-scale purification is generally not enough for initial crystallization trials, even with low-volume crystallization, and thus scaled-up protein production and purification is needed. The purification tags can be removed upon large-scale purification.

4.4.4.1. Autoinduction

Protein expression under control of the T7 *lac* promoter system can be induced either with the chemical inducer isopropyl-β-galactoside (IPTG) or by autoinduction using a mixture of glucose, glycerol and lactose during *E. coli* growth. In autoinduction, cells grow to relatively high density in a defined ratio of glucose-to-lactose media (Studier, 2005). Initially, glucose prevents induction by lactose. When available glucose is used up, lactose then induces target protein expression switching to lactose metabolism.

Cell cultures (1 ml) are grown in 2.2 ml deep-well 96-well blocks. Protein expression is autoinduced using commercially available glucose/lactose media (*e.g.* Overnight Express from EMD Biosciences). The following day, cells are lysed either by repeated freezing and thawing cycles, or by addition of a lysozyme solution. A combination of lysozyme and benzonase solutions (*e.g.* PopCulture reagent from Novagen) eliminates the need for cell harvesting prior to lysis and lysate clarification following lysis. The cell lysate can then be directly used for expression and purification tests. Alternatively, whole-cell lysates can be filtered through a 96-well filter plate, allowing for separation of inclusion bodies from the soluble fraction (filtrate) so that protein expression in the soluble fraction can be assessed.

4.4.4.2. Expression and solubility test: dot blot

Dot blot is a simple method that can be used to analyse either total protein expression in the cell lysate, or soluble protein expression following the separation of supernatant and cell pellet. Protein samples from the total cell lysate in a 96-well plate are dotted onto a nitrocellulose membrane by applying a vacuum. Target proteins are then probed with an antibody against the

protein or against an affinity tag, *e.g.* anti-His-tag antibody for His-tagged protein detection (Fig. 4.4.3.1c). Alternatively, the cell lysates are analysed by sodium dodecyl sulfate–polyacrylamide gel electrophoresis (SDS–PAGE) for protein expression using a pre-cast 48- or 96-well SDS gel (E–PAGE 48 or 96 from Invitrogen). The gel consists of 48 or 96 wells for samples, and an additional four or eight wells for protein markers. The protein can be loaded either with a liquid-handling robot or with a multi-channel pipettor, and electrophoresis is completed within 15 min (Fig. 4.4.3.1d).

4.4.4.3. Small-scale purification

Affinity purifications are preferred in a 96-well format because the specific interaction between the protein and affinity resin allows for a simple ‘bind–wash–elution’ procedure. Purification using a His-tag or a GST-tag (where GST = glutathione S-transferase) is popular in structure determination projects, and appropriate resins are available as magnetic beads or agarose resin in a 96-well plate, *i.e.* magnetic Ni-NTA or GST beads, or Ni- or Co-linked agarose discs, respectively.

The soluble fraction or whole-cell lysate is transferred to a 96-well plate containing an affinity resin for protein binding. The beads are separated from unbound protein by placing them in a magnetic stand designed to accommodate a 96-well format, or by filtration using a 96-well filter plate that retains the beads but allows passage of the cell lysate. Beads are washed, and bound protein is then eluted with the appropriate elution buffer (*i.e.* imidazole and reduced glutathione for His-tagged and GST-tagged proteins, respectively) into a 96-well plate by applying a vacuum (either as part of an appropriately equipped liquid-handling robot, or by using a 96-well vacuum manifold) or by centrifugation. Purified proteins are analysed by SDS–PAGE using a pre-cast 48- or 96-well protein gel. The gels can then be stained either by protein staining or western blotting. A new system for protein and DNA analysis has been developed based on a microfluidic chip (Caliper Labchip, Caliper Life Science). These chips contain a network of miniaturized channels, through which fluids and chemicals are moved to separate DNAs or proteins. The DNA and protein signal is measured by laser-induced fluorescence. The instrument is capable of separating 2 ng–2 µg amounts of protein, one well after another; it takes about 1 h to analyse a 96-well plate.

4.4.5. Crystallization

HT crystallization using 96-well crystallization plates greatly reduces the total amount of protein required for screening and thus structure determination. Robotic crystallization systems are capable of dispensing nanolitre droplets (<100 nl) and hence substantially increase the number of conditions that can be screened with a fixed amount of protein sample, as well as reducing the time required for setting up a series of crystallization trials. A thousand conditions with different crystallization parameters (*e.g.* pH, salts, temperature) can be screened with 100 µl of protein sample. At the time of writing, commercially available crystallization robots include the Honey Bee (Cartesian Micro-array), Phoenix (Rigaku), Mosquito (Molecular Dimensions Limited) and Oryx (Douglas Instrumentation). These crystallization robots can set up either vapour-diffusion, microbatch or hanging-drop methods in 96-well plates within 15 min. Crystallization robots can also be integrated into a larger

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system that includes storage and imaging of crystallization plates, and liquid-formulation robots.

Honey Bee and Phoenix robots contain a single (or several) non-contact channel that dispenses protein solution and a 96-channel dispenser head that dispenses the crystallization solutions (50 nl to 100 μ l). The 96-channel head transfers crystallization solutions from a 96-well deep-well plate into the reservoir and crystallization drop in a 96-well crystallization plate (Fig. 4.4.3.1e). A single channel transfers protein solution into each of the 96-well drops one by one without touching the precipitate drops. The plate is then sealed with a clear film by the user. The Mosquito liquid-handling robot can set up drops with a hanging-drop geometry, and is more popular with membrane protein crystallization (detergent solutions have a tendency to adhere to the side of the drop well in sitting-drop geometry). The Mosquito uses only disposable tips capable of dispensing 20 nl–1.2 μ l volume; thus the user can control the location of drop deposition more precisely because the disposable tips can touch the drop. The use of disposable tips also prevents cross-contamination between samples, and washing steps between samples are eliminated.

The Fluidigm TOPAZ system utilizes a new technology, the crystallization screen chip, in which protein sample and reagent solutions are automatically loaded into diffusion chambers within the protein screen chip and the two solutions mixed by free interface diffusion, as opposed to vapour diffusion or microbatch techniques (Thorsen *et al.*, 2002). A very small amount of protein is required for a crystallization screen, *i.e.* as little as 1.0 μ l protein solution for 96 trials. Crystals obtained from the protein chip are generally too small for X-ray data collection, and thus need to be scaled up to obtain diffraction-quality crystals.

Progress of crystallization trials from large numbers of 96-well plates can be monitored using an imaging robot to take pictures of individual crystallization drops. The resulting images can then be analysed either manually or using automatic crystal recognition systems at specified time intervals (Markley *et al.*, 2009). Remote viewing of recorded crystal pictures is also available over the web. Each recorded image is linked to crystallization conditions for evaluation and scoring of the crystallization conditions (Fig. 4.4.3.1f). Minstrel (Rigaku), CrystalFarm (Bruker) or HomeBase (The Automation Partnership) systems offer integrated systems for plate storage and imaging.

Crystallization conditions that initially produced crystals should be optimized to improve crystal growth and quality. A liquid-handling robot can be used to make screens in 96-well deep-well plates. A liquid-formulation robot has been developed for protein crystallization to make grid screens of 96-well deep-well conditions (*e.g.* Alchemist from Rigaku). The crystallization conditions stored in an imaging robot are linked to the liquid-formulation software, and can be used to formulate 96-well screen conditions for optimization experiments. Other optimization methods such as crystallization in gels, control of nucleation using oil mixtures or microporous materials, and seeding experiments can also be employed in an HT fashion (Chayen, 2003; Georgiev *et al.*, 2006; Sugahara *et al.*, 2008).

4.4.6. Synchrotron data collection

High-brilliance beamlines at modern synchrotrons have significantly reduced the time required for X-ray data collection, and complete data sets can often be collected within minutes. Thus, the time required for crystal mounting and centring is no longer

negligible. Automatic crystal mounting and centring allow users to remotely mount crystals for crystal evaluation and data collection without entering the experimental hutch (Manjasetty *et al.*, 2008; Sharff & Jhoti, 2003; Sugahara *et al.*, 2008).

Automated crystal mounting allows the screening of many crystals for diffraction quality and then goes back to the best diffraction-quality crystals for full data collection. The mounting robot picks up frozen crystals in a pin from a Dewar, puts them on a goniometer and retrieves the pin after the diffraction test. The automated crystal-mounting robots have primarily been developed for use at synchrotron sources, although a commercial version of an automatic sample-mounting robot is now available that can be used with a home-source X-ray generator (ACTOR from Rigaku and cryogenic sample changer from Marresearch). Fully automated crystal alignment is not yet available, but semi-automated crystal centring by clicking a mouse to indicate the intended centre of the crystal is used at most beamlines.

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