XACT-seq: A photocrosslinking-based technique for detection of the RNA polymerase active-center position relative to DNA in *Escherichia coli*

XACT-seq ("crosslink between active center and template sequencing") is a technique for high-throughput, single-nucleotide resolution mapping of RNA polymerase (RNAP) active center positions relative to the DNA template. XACT-seq overcomes limitations of approaches that rely on analysis of the RNA 3’ end (e.g., native elongating transcript sequencing) or that report RNAP positions with low resolution (e.g., ChIP-seq and ChIP-exo). XACT-seq can be used to map RNAP active center positions in transcription initiation complexes, initially transcribing complexes, and transcription elongation complexes.
Protocol

XACT-seq: A photocrosslinking-based technique for detection of the RNA polymerase active-center position relative to DNA in *Escherichia coli*

Chirangini Pukhrambam, 1,2 Irina O. Vvedenskaya, 1,2 and Bryce E. Nickels 1,3,*

1Department of Genetics and Waksman Institute, Rutgers University, Piscataway, NJ 08854, USA
2Technical contact
3Lead contact
*Correspondence: bnickels@waksman.rutgers.edu
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SUMMARY

XACT-seq (“crosslink between active-center and template sequencing”) is a technique for high-throughput, single-nucleotide resolution mapping of RNA polymerase (RNAP) active-center positions relative to the DNA template. XACT-seq overcomes limitations of approaches that rely on analysis of the RNA 3′ end (e.g., native elongating transcript sequencing) or that report RNAP positions with low resolution (e.g., ChIP-seq and ChIP-exo). XACT-seq can be used to map RNAP active-center positions in transcription initiation complexes, initially transcribing complexes, and transcription elongation complexes.

For complete details on the use and execution of this protocol, please refer to Winkelman et al. (2020).

BEFORE YOU BEGIN

**Introduction**

XACT-seq involves formation of transcription complexes using an RNAP derivative that has the photo-activatable crosslinking amino acid p-benzoyl-L-phenylalanine (Bpa) (Chin et al., 2002) incorporated at RNAP β subunit residue R1148 (RNAP-β<sup>R1148Bpa</sup>) which, upon photo activation, forms covalent crosslinks with DNA at a position exactly 5 nt downstream of the RNAP active-center A site (Figures 1A and 1B) (Yu et al., 2017). XACT-seq is performed using merodiploid *Escherichia coli* cells containing a plasmid-encoded, decahistidine-tagged RNA polymerase (RNAP) derivative and a chromosomally encoded, untagged RNAP (Figure 1C). In this case, plasmid, pIA900-β<sup>R1148Bpa</sup>, contains a gene for the RNAP β′ subunit with a nonsense codon (TAG) at position 1148 and a decahistidine coding sequence (Figure 1C). A second plasmid, pEVOL-pBpF, contains a gene for an engineered Bpa specific UAG suppressor tRNA and a gene for an engineered Bpa specific aminoacyl tRNA synthetase (Figure 1C). Growth of cells containing both plasmids in media supplemented with Bpa results in production of β′<sup>R1148Bpa</sup> and corresponding formation of transcription complexes containing RNAP-β<sup>R1148Bpa</sup> (Figures 1B and 1C).

To initiate covalent crosslinking of RNAP to DNA, cells are UV irradiated (Figure 2, step 1). After UV irradiation, cells are lysed, RNAP-DNA complexes are recovered by immobilized metal ion affinity chromatography (IMAC) targeting the decahistidine tag on RNAP-β<sup>R1148Bpa</sup> (Figure 2, step 2), and crosslink positions and crosslink yields are defined by high-throughput sequencing of primer extension products (Figure 2, steps 3–6). The procedure enables the position of the RNAP active-center relative to DNA to be mapped, in vivo, with single-nucleotide resolution.

Below we describe use of XACT-seq to analyze transcription complexes associated with up to at least ~4 million promoter sequences in *E. coli* (Figure 3). In this case, XACT-seq is performed with cells
containing a plasmid-borne library of barcoded template sequences (e.g., placCONS-N11, see Figure 3). For the details on construction of placCONS-N11 library see (Vvedenskaya et al., 2015, 2018; Winkelman et al., 2020).

Prepare growth media and buffers

© Timing: 1 day

1. Lysogeny Broth (LB): dissolve components in 1 L of ddH₂O and autoclave. Store at 25°C.
2. LB Agar: dissolve components in 1 L of ddH₂O and autoclave. Pour ~25 mL of media into each 100 x 15 mm Petri dish. Store at 4°C.
3. **Resuspension Buffer**: combine components in 0.9 L of ddH₂O, adjust pH to 8 and add ddH₂O to 1 L. Filter using a sterile vacuum filter unit (1 L, 0.2 μm pore size). Store at 25°C.

| Reagent               | Final concentration | Amount/volume |
|-----------------------|---------------------|---------------|
| Na₂HPO₄ · 7H₂O        | 0.05 M              | 13.4 g        |
| NaCl                  | 1.4 M               | 81.9 g        |
| Imidazole (C₃H₄N₂)   | 0.02 M              | 1.36 g        |
| Tween 20              | 0.1%                | 1 mL          |
| Ethanol (100%)        | 5%                  | 50 mL         |

**Note**: immediately before use, add β-mercaptoethanol (BME) to a concentration of 15 mM.

4. **Ni-NTA Wash Buffer**: combine components in 0.9 L of ddH₂O, adjust pH to 8 and add ddH₂O to 1 L. Filter using a sterile vacuum filter unit (1 L, 0.2 μm pore size). Store at 4°C.

| Reagent               | Final concentration | Amount/volume |
|-----------------------|---------------------|---------------|
| Na₂HPO₄ · 7H₂O        | 50 mM               | 13.4 g        |
| NaCl                  | 300 mM              | 17.5 g        |
| Imidazole (C₃H₄N₂)   | 30 mM               | 2.0 g         |
| Tween 20              | 0.1%                | 1 mL          |
| Ethanol (100%)        | 5%                  | 50 mL         |

**Note**: immediately before use, add BME to a concentration of 15 mM.

5. **Ni-NTA Elution Buffer**: combine components in 0.9 L of ddH₂O, adjust pH to 8 and add ddH₂O to 1 L. Filter using a sterile vacuum filter unit (1 L, 0.2 μm pore size). Store at 4°C.

| Reagent               | Final concentration | Amount/volume |
|-----------------------|---------------------|---------------|
| Na₂HPO₄ · 7H₂O        | 50 mM               | 13.4 g        |
| NaCl                  | 300 mM              | 17.5 g        |
| Imidazole (C₃H₄N₂)   | 300 mM              | 20.4 g        |
| Tween 20              | 0.1%                | 1 mL          |
| 100% Ethanol          | 5%                  | 50 mL         |

**Note**: immediately before use, add BME to a concentration of 15 mM.

6. **Storage Buffer (2×)**: combine components in 80 mL of ddH₂O, adjust pH to 8 and add ddH₂O to 100 mL. Filter using a sterile vacuum filter unit (0.2 μm pore size). Store at 4°C.

| Reagent               | Final concentration | Volume |
|-----------------------|---------------------|--------|
| Tris-Cl pH 8.0 (1 M)  | 20 mM               | 2 mL   |
| KCl (2 M)             | 200 mM              | 10 mL  |
| MgCl₂ (1 M)           | 20 mM               | 2 mL   |
| EDTA (0.5 M)          | 0.2 mM              | 0.04 mL|
| DTT (1 M)             | 1 mM                | 0.1 mL |

**Note**: immediately before use, add BME to a concentration of 15 mM.
7. TBE (10×): dissolve components in 1 L of ddH₂O. Store at 25°C.

| Reagent                        | Final concentration | Amount  |
|--------------------------------|---------------------|---------|
| Tris base                      | 20 mM               | 108 g   |
| Boric Acid                     | 200 mM              | 55 g    |
| EDTA, disodium salt, dihydrate | 20 mM               | 7.4 g   |

maximum time for storage: 4 months

**KEY RESOURCES TABLE**

| REAGENT or RESOURCE | SOURCE                          | IDENTIFIER |
|---------------------|---------------------------------|------------|
| **Bacterial strains** |                                 |            |
| NiCo21 (DE3)        | NEB                             | Cat#C2529H |
| **Chemicals, peptides, and recombinant proteins** |                                 |            |
| Nuclease-Free Water (not DEPC-treated) | Thermo Fisher Scientific | Cat#A09932 |
| Bacto agar          | VWR                             | Cat#90000-760 |
| Bacto tryptone      | VWR                             | Cat#90000-286 |
| Bacto yeast extract | VWR                             | Cat#90000-726 |
| Chloramphenicol     | Gold Biotech                    | Cat#C-105-25 |
| Spectinomycin       | Duchefa Biochemie               | Cat#S0188.0025 |
| Streptomycin        | Thermo Fisher Scientific        | Cat#15140122 |
| Carbenicillin       | Gold Biotech                    | Cat#C-103-25 |
| Rifampicin          | Gold Biotech                    | Cat#R-120-25 |
| H-Bpa-OH            | Bachem                          | Cat#F-2800 |
| IPTG Gold           | Biotech                         | Cat#H481C50 |
| SOC Outgrowth Medium| NEB                             | Cat#B9020S |
| Lysozyme Egg White  | Gold Biotech                    | Cat#L-040-10 |
| Glycerol, nuclease free| VWR                            | Cat#EM-4750 |
| Tris base (Amresco) | VWR                             | Cat#97061-800 |
| Boric acid (ACS grade)| VWR                            | Cat#97061-980 |
| EDTA disodium salt dihydrate | VWR            | Cat#97061-018 |
| Imidazole           | VWR                             | Cat#EM-5720 |
| Formamide, deionized | VWR                            | Cat#EM-4610 |
| Sodium dodecylsulfate (SDS)| VWR              | Cat#97064-470 |
| Bromophenol blue    | VWR                             | Cat#EM-BX1410-7 |
| Xylene Cyanol       | Sigma-Aldrich                   | Cat#X4126-10G |
| 0.5 M EDTA pH 8     | Thermo Fisher Scientific        | Cat#AM9260G |
| 3 M Sodium Acetate pH 5.5 | Thermo Fisher Scientific      | Cat#AM9740 |
| BSA, Molecular Biology Grade | NEB                      | Cat#B90005 |
| 5 M Betaine solution| VWR                             | Cat#101375-612 |
| Dimethyl sulfoxide  | VWR                             | Cat#BDH1115-1LP |
| Heparin sulfate     | Sigma-Aldrich                   | Cat#H-3393 |
| Glycogen from oyster (type II) | Sigma-Aldrich             | Cat#GB751 |
| Ethyl alcohol       | Pharmco-AAPER                   | Cat#111000200 |
| Isopropl alcohol    | VWR                             | Cat#BDH1133-1LP |
| Low Range ssRNA Ladder | NEB                       | Cat#N0364S |
| O’Gene Ruler Ultra Low Range DNA Ladder | Thermo Fisher Scientific  | Cat#SM1223 |
| 6x Orange DNA Loading Dye | Thermo Fisher Scientific    | Cat#R0631 |
| SYBR Gold Nucleic Acid Gel Stain | Thermo Fisher Scientific | Cat#S11494 |
| Phenol:Chloroform:IAA pH 8 | Thermo Fisher Scientific | Cat#AM9732 |
| Taq DNA polymerase  | NEB                             | Cat#M0273 |
| 5’ App DNA/RNA ligase | NEB                        | Cat#M03195 |
| T4 RNA Ligase 1 (sRNA Ligase) | NEB                   | Cat#M0204L |

(Continued on next page)
STEP-BY-STEP METHOD DETAILS

Introduction of plasmids into E. coli cells

- **Timing:** ~18 h for step 1
- **Timing:** ~5 h for step 2
- **Timing:** ~18 h for step 3
Figure 3. Three plasmid system to apply XACT-seq to a promoter library of up to at least ~4 million sequences

Top shows cells containing decahistidine-tagged RNAP βR1148Bpa and a promoter library of ~4 million sequences (plasmid placCONS-N11). Bottom shows sequence of the placCONS-N11 promoter library. Blue, promoter -35 and -10 elements; dark yellow, randomized sequences from position +3 to +13; light yellow, transcribed region barcode sequence. Other symbols and colors as in Figure 1.

© Timing: ~5 h for step 4

© Timing: ~18 h for step 5

Note: We have found that the sequential introduction of plasmid library placCONS-N11, followed by plasmid pIA900-βR1148Bpa, followed by plasmid pEVOL-pBpF into E. coli cells provides consistent results.

1. First step to generate three-plasmid merodiploid system to analyze up to ~4 million promoter sequences: introduction of placCONS-N11 library into E. coli.

Note: Because of the total number of sequences present in the placCONS-N11 library (up to ~4 million promoter sequences) our lab uses electrocompetent cells for each transformation step. In general, use of electrocompetent cells results in a higher transformation efficiency compared with use of chemically competent cells.

a. Combine 50 μL of NiCo21 (DE3) electrocompetent cells with 1 μL (90–100 ng) of placCONS-N11 in a microcentrifuge tube.
b. Transfer the mixture to a 1 mm electroporation cuvette and apply current using an electroporator. (Our lab uses a MicroPulser electroporator and a pulse time of 3.5–4 ms.)
c. Add 950 μL of SOC (prewarmed to 37°C), mix by pipetting, and transfer to a 17 mm × 100 mm round-bottom culture tube.
d. Incubate for 1 h at 37°C with gentle mixing. (Our lab uses a Cel-Gro Tissue Culture Rotator Drum set to ~55 rpm.)
e. Remove cells from culture tube and mix with 50 mL of LB containing 50 μg/mL streptomycin and 50 μg/mL spectinomycin in a 250 mL flask.
f. Shake flask in orbital shaker at 220 rpm for 16 h at 37°C.

**Note:** we recommend performing steps 1.a-b in a cold room.

**Note:** The desired number of transformants is \( \sim 5 \times 10^6 \). To estimate the number of transformants, a serial dilution is prepared using the cell suspension prior to step 1.e. Each dilution is plated on LB agar containing 50 µg/mL streptomycin and 50 µg/mL spectinomycin and is grown for 16 h at 37°C. The number of colonies on each plate is counted. The number of transformants is calculated assuming each colony is derived from an independent transformation event. For additional details see: Troubleshooting, Problem 1.

2. Next, electrocompetent cells of NiCo21 (DE3) containing the placCONS-N11 library are generated prior to introduction of plA900-\( \beta^{R1148Bpa} \).

   a. In a 500 mL flask inoculate 100 mL of LB medium containing 50 µg/mL streptomycin and 50 µg/mL spectinomycin with 1 mL of the cell suspension from step 1.f.
   b. Place flask on orbital shaker set to 220 rpm and incubate at 37°C until the cell suspension reaches an OD\(_{600}\) of 0.5 (~2 h).
   c. Transfer cell suspensions into two 50-mL centrifuge tubes submerged in ice.
   d. Collect cells by centrifugation (3000 \( \times \) g; 4°C; 10 min), remove the supernatant and resuspend cells in 50 mL of ddH\(_2\)O kept at 4°C.
   e. Repeat step 2.d two more times.
   f. Resuspend cells in 10 mL of 10% glycerol kept at 4°C. Collect cells by centrifugation (3000 \( \times \) g; 4°C; 10 min) and remove the supernatant.
   g. Repeat step 2.f two more times and resuspend cells in 0.5 mL of 10% glycerol kept at 4°C.

   **Note:** Do not vortex tubes while resuspending cell pellets at wash steps 2.d-g. Instead, swirl the tubes until cells are resuspended. In addition, if you decide to transfer cell suspensions into new tubes at step 2.g, use wide bore pipette tips.

   **Note:** We recommend using freshly prepared competent cells for the next transformation step, otherwise cells can be stored at \(-80°C\) for up to 6 months.

3. Second step to generate three-plasmid merodiploid system to analyze up to \( \sim 4 \) million promoter sequences: introduction of plasmid plA900-\( \beta^{R1148Bpa} \) into NiCo21 (DE3) cells containing the placCONS-N11 library.

   a. Combine 50 µL of NiCo21 (DE3) cells containing the placCONS-N11 library (prepared in step 2.g above) with 1 µL (100–120 ng) of plA900-\( \beta^{R1148Bpa} \) in a microcentrifuge tube. Transfer the mixture to a 1 mm electroporation cuvette and apply current using an electroporator. (Our lab uses a MicroPulser electroporator and a pulse time of 3.5–4 ms.)
   b. Add 950 µL of SOC (prewarmed to 37°C), mix by pipetting, and transfer to a 17 mm \( \times \) 100 mm round-bottom culture tube.
   c. Incubate for 1 h at 37°C in Cel-Gro Tissue Culture Rotator Drum set to \( \sim 55 \) rpm.
   d. Remove cells from culture tube and mix with 50 mL of LB containing 50 µg/mL streptomycin, 50 µg/mL spectinomycin and 100 µg/mL carbenicillin in a 250 mL flask.
   e. Place flask on orbital shaker set to 220 rpm for 16 h at 37°C.

   **Note:** We recommend performing step 3.a-b in a cold room.

   **Note:** The desired number of transformants is \( \sim 5 \times 10^6 \). To estimate the number of transformants, prepare a serial dilution of the cell suspension prior to step 3.d. Plate each dilution on LB agar containing 50 µg/mL streptomycin, 50 µg/mL spectinomycin and 100 µg/mL carbenicillin. Incubate plates for 16 h at 37°C. Count the number of colonies on each plate. Calculate the
number of transformants assuming each colony is derived from an independent transformation event. For additional details see: Troubleshooting, Problem 1.

4. Next, electrocompetent cells of NiCo21 (DE3) containing pIA900-pR1148Bpa and the placCONS-N11 library are prepared prior to introduction of pEVOL-pBpF.
   a. Inoculate 100 mL of LB medium containing 50 μg/mL streptomycin, 50 μg/mL spectinomycin and 100 μg/mL carbenicillin in 500 mL flask with 1 mL of the cell suspension from step 3.e.
   b. Place flask in orbital shaker set to 220 rpm and incubate at 37°C until the cell suspension reaches an OD₆₀₀ of 0.5 (~2 h).
   c. Transfer cell suspensions into two 50 mL centrifuge tubes submerged in ice.
   d. Collect cells by centrifugation (3000 × g; 4°C; 10 min) and remove the supernatant. Resuspend cells in 50 mL of ddH₂O kept at 4°C.
   e. Perform step 4.d two more times.
   f. Resuspend cells in 10 mL of 10% glycerol kept at 4°C by pipetting. Collect cells by centrifugation (3000 × g; 4°C; 10 min) and remove the supernatant.
   g. Repeat step 4.f two more times and resuspend cells in 0.5 mL of 10% glycerol kept at 4°C.

   **Note:** Do not vortex tubes while resuspending cell pellets at wash steps 4.d-g. Instead, swirl the tubes until cells are resuspended. In addition, if you decide to transfer cell suspensions into new tubes at step 4.g, use wide bore pipette tips.

   **Note:** We recommend using freshly prepared competent cells for the next transformation step, otherwise cells can be stored at −80°C for up to 6 months.

   ### Pause point: Cells can be stored at −80°C for up to 6 months.

5. Third step to generate three-plasmid merodiploid system to analyze up to ~4 million promoter sequences: introduction of plasmid pEVOL-pBpF into NiCo21 (DE3) cells containing pIA900-pR1148Bpa and the placCONS-N11 library.
   a. Combine 50 μL of NiCo21 (DE3) cells containing pIA900-pR1148Bpa and the placCONS-N11 library (prepared in step 4.g) with 1 μL (~8 ng) of pEVOL-pBpF in a microcentrifuge tube. Transfer the mixture to a 1 mm electroporation cuvette and apply current using an electroporator. (Our lab uses a MicroPulser electroporator and a pulse time of 3.5–4 ms.)
   b. Add 950 μL of SOC (prewarmed to 37°C), mix by pipetting, and transfer to a 17 mm × 100 mm round-bottom culture tube.
   c. Incubate for 1 h at 37°C with gentle mixing. (Our lab uses a Cel-Gro Tissue Culture Rotator Drum set to ~55 rpm.)
   d. Centrifuge cell suspension to pellet cells (21,000 × g; 25°C; 1 min) and remove 800 μL media. Resuspend cells in the remaining 200 μL media. Plate 100 μL of the cell suspension on two LB agar 100 × 15 mm Petri dishes containing 50 μg/mL spectinomycin, 50 μg/mL streptomycin, 100 μg/mL carbenicillin, and 25 μg/mL chloramphenicol.
   e. Incubate plates at 37°C for 16 h.

   **Note:** Perform step 5.a-b in a cold room.

   **Note:** Each LB agar plate will contain a lawn of transformants. We estimate the number of transformants as 11–16 × 10⁶ per plate.

**Growth and UV irradiation of plasmid-containing cells**

© Timing: ~6–8 h
6. Expression of Bpa-labeled RNAP in NiCo21 (DE3) cells containing pEVL-pBpF, pIA900-R1148Bpa, and the placCONS-N11 library followed by UV-irradiating cells to induce crosslinking between Bpa-labeled RNAP and DNA.

a. Prepare 2.5 mL of 100 mM Bpa by dissolving 67.5 mg Bpa in 2.5 mL of 1 M NaOH.

b. Prepare 250 mL LB media containing 1 mM Bpa by adding the 100 mM Bpa solution prepared in step 6.a dropwise into 250 mL LB media while stirring. Adjust the pH to 7.2 with 1 M HCl (add ~ 0.8 mL of 1 M HCl per 100 mL LB media).

c. Add 100 µg/mL carbenicillin, 50 µg/mL spectinomycin, 50 µg/mL streptomycin and 25 µg/mL chloramphenicol to the media prepared in step 6.b.

**Note:** Bpa is a photoreactive compound. Therefore, steps 6.a-b should be performed in a room with minimal light and performed in as short a time period as possible.

d. To recover transformants from step 5.e, add 7 mL of LB to the surface of each agar plate and resuspend cells using a 6” sterilized wooden applicator. Combine the cell suspensions from each plate in a 50 mL Falcon tube and measure the OD600 of the mixture.

**Note:** We recommend making a 1:10 dilution of cell suspension for accurate measurement of OD600.

e. Using the mixture prepared in step 6.d, inoculate 250 mL LB containing 1 mM Bpa, 100 µg/mL carbenicillin, 50 µg/mL spectinomycin, 50 µg/mL streptomycin and 25 µg/mL chloramphenicol to an OD600 of ~0.3. (For example, if the OD600 of the undiluted cell suspension is 7.5, then adding ~10 mL of this suspension to 250 mL media will yield an OD600 of ~0.3).

f. Place the mixture in a 1 L flask in an orbital shaker set to 220 rpm and incubate for 1 h at 37°C.

**CRITICAL:** Bpa is a photoreactive compound. Therefore, the exposure of the cultures to light during growth should be minimized by covering culture flasks with aluminum foil and incubating cells in an unlit room or growth chamber.

g. Add IPTG to a final concentration of 1 mM. Incubate cultures for additional 3 h at 37°C (220 rpm).

h. Transfer 9 mL of the cell suspension into a 13 mm x 100 mm borosilicate glass test tubes (total of 27 tubes per sample).

i. UV-irradiate cell suspensions. To ensure reproducibility, we recommend use of a commercially available photoreactor. Our lab uses a Rayonet RPR-100 photochemical reactor (350 nm wavelength, 20 min, 25°C).

j. Transfer cell suspensions from tubes to 250-mL centrifugal bottles. Collect cells by centrifugation (3000 × g; 4°C; 10 min), remove supernatant, and store cell pellets at −20°C.

**Note:** To trap static initial-transcribing complexes containing 2- to 3-nt RNA products in vivo, despite the presence of all NTP substrates in vivo, the RNAP inhibitor rifampin (Rif), which blocks extension of RNA products beyond a length of 2–3 nt, can be added to a final concentration of 200 µg/mL prior to UV irradiation. After adding rifampin, we recommend incubating cells for 10 min at 37°C (220 rpm).

**Pause point:** Cell pellets can be stored at −20°C for 2–3 weeks.

### Isolation of crosslinked complexes from cell lysates

- **Timing:** ~4–6 h for step 7
- **Timing:** ~1 h for step 8
7. Use of immobilized metal-ion affinity chromatography (IMAC) to isolate Bpa-labeled RNAP from cell lysate
   a. Thaw frozen cell pellets prepared in step 6.j at 4°C for 30 min and resuspend in 40 mL of Resuspension Buffer containing 15 mM BME and 2 mg lysozyme.
   b. Sonicate cells at 4°C (Our lab uses a Sonics Vibra Cell instrument set at 10 s pulse/10 s pause for 5 min).
   c. Remove cell debris by centrifugation (23,000 × g; 30 min; 4°C) to form compact pellet of debris.
   d. Aliquot 1 mL of Ni-NTA agarose in 50-mL Falcon tube. Wash the resin with 1 mL of nuclease-free water and equilibrate in 1 mL of Resuspension Buffer. Repeat the equilibration step two more times.
   e. Add supernatant from step 7.c to the resin. Incubate for 30 min at 4°C with gentle rocking on rotator (Our lab normally uses a Stovall Belly Dancer Shaker). Keep tube on ice and cover with foil to protect from light.
   f. Load the slurry into a 15 mL polyprep column to collect the Ni-NTA-agarose resin.
   g. Wash resin with 10 mL of 1 X Ni-NTA Wash Buffer.
   h. Add 3 mL of 1 X Ni-NTA Elution Buffer containing 300 mM imidazole, incubate 1 min and elute His-tagged proteins from the resin.
   i. Concentrate the eluate to ~0.1–0.2 mL by centrifugation (3000 × g; 7–12 min, 4°C) using 100K MWCO Amicon Ultra-4 centrifugal filter.
   j. To perform buffer exchange, add 2 mL of 2 X Storage Buffer to the concentrated eluate. Reduce the volume of the mixture to ~0.1–0.2 mL by centrifugation (3000 × g; 7–12 min, 4°C).
   k. Repeat step 7.j.
   l. Add an equal volume of 100% glycerol (~0.1–0.2 mL) to the concentrated eluate. Mix thoroughly by pipetting and store sample at −80°C.

△ CRITICAL: Perform steps 7.a-l at 4°C.

Note: The eluate at step 7.l contains Bpa-labeled RNAP that is crosslinked to DNA and Bpa-labeled RNAP that is not crosslinked to DNA.

Note: After step 7.l, the concentration of protein in the eluate is typically ~3.5–5 μM.

Pause point: The eluate can be stored at −80°C for at least 3 months.

8. Heat denaturation of crosslinked complexes and binding to MagneHis Ni-particles
   The next step of the protocol involves heat denaturation of crosslinked complexes followed by binding to MagneHis Ni-particles. This step removes non-crosslinked DNA from the reaction mixture.
   a. Aliquot 20 μL MagneHis Ni-particles in two 1.5 mL tubes. Remove storage solution and equilibrate the particles in 50 μL of 1 X Taq DNA polymerase buffer using MagneSphere magnetic separation stand (Promega). Repeat equilibration step. Keep tubes on ice while preparing the crosslinked complexes for binding with MagneHis Ni-particles.
   b. Set up two tubes containing 25 μL of the eluate prepared in step 7.l. Add 25 μL of nuclease-free H2O, 12 μL of 5 M NaCl, and 0.5 μL of 1 mg/mL heparin to each tube.
   c. Incubate the mixtures at 95°C for 5 min to denature dsDNA. Cool to 4°C and incubate for 5 min.
   d. Add the denatured mixture to MagneHis Ni-particles from step 8.a.
   e. To bind the crosslinked complexes to beads, incubate for 10 min at 25°C. Gently flick the tube 2–3 times every 2 min to mix the sample.
   f. Collect the beads using a magnetic separation stand. Discard the supernatant.
g. Add 50 µL of wash buffer containing 10 mM Tris-Cl pH 8.0, 1.2 M NaCl, 10 mM MgCl₂ and 10 µg/mL heparin. Incubate 10 min at 25°C. Gently flick the tube 2–3 times every 2 min to mix the sample.

h. Collect the beads using a magnetic separation stand. Discard the supernatant.

i. Add 50 µL of 1 X Taq DNA polymerase buffer (NEB) to wash the beads and to prepare samples for the next step. Gently flick the tube 2–3 times to mix the sample.

j. Collect the beads using a magnetic separation stand. Discard the supernatant.

k. Repeat steps 8.i-j.

l. Resuspend the beads in 10 µL of 1 X Taq DNA polymerase buffer.

m. Pool the beads in one tube (total volume ~20 µL). Store at 4°C while preparing primer extension reactions.

△ CRITICAL: Proceed immediately to primer extension step.

**Primer extension reactions and purification of primer extension products**

⊙ Timing: ~4 h for step 9

⊙ Timing: ~18 h for step 10

⊙ Timing: ~20 h for step 11

The next step of the protocol involves primer extension reactions using oligonucleotide complementary to sequences downstream of the N11 region of the pLacCONS-N11 library.

9. Primer extension reaction
   a. Prepare primer extension reaction:

| Reagent                                          | Final concentration | Amount |
|--------------------------------------------------|---------------------|--------|
| 10 X Taq DNA Polymerase Buffer                   | 1 X                 | 10 µL  |
| s128A, Primer Extension oligo (10 µM)            | 0.2 µM              | 2 µL   |
| Taq DNA Polymerase (5 U/µL)                      | 0.1 U/µL            | 2 µL   |
| 10 X dNTPs (2.5 mM dATP, 2.5 mM dCTP, 2.5 mM dGTP, 2.5 mM dTTP) | 0.25 mM each        | 10 µL  |
| 5 M Betaine                                      | 2 M                 | 40 µL  |
| DMSO (100%)                                      | 5%                  | 5 µL   |
| Nuclease-free H₂O                                | n/a                 | 15 µL  |
| Crosslinked complexes on MagneHis beads          | n/a                 | 16 µL  |
| Total                                            | n/a                 | 100 µL |

b. Split each reaction in two 0.2 mL PCR tubes (50 µL reaction mixture in each tube) and place in a thermocycler.

c. Perform primer extension reactions using the following thermocycler conditions:

| Steps                | Temperature | Time | Cycles |
|----------------------|-------------|------|--------|
| Initial Denaturation | 95°C        | 30 s | 1      |
| Denaturation         | 95°C        | 30 s | 40 cycles |
| Annealing            | 55°C        | 30 s |        |
| Extension            | 72°C        | 30 s |        |
| Final Extension      | 72°C        | 5 min| 1      |
| Hold                 | 4°C         |      | forever|

△ CRITICAL: Proceed immediately to primer extension step.
d. Pool 2 reactions into a single 1.7 mL tube.

Note: We recommend optimizing the number of cycles in primer extension reaction (i) to generate enough products for downstream steps in the protocol and (ii) to avoid non-specific background products.

10. Purification of primer extension products using phenol/chloroform/IAA mixture
Next, purification of primer extension products is performed using phenol/chloroform/IAA mixture followed by ethanol precipitation.

a. Add 100 µL phenol:chloroform:IAA (pH 8.0), mix by vortexing for 30 s, centrifuge (21,000 × g; 1 min; 25°C), and recover the upper aqueous phase (~100 µL).

b. Add 10 µL of 3M NaOAc, 1.1 µL of 10 mg/mL glycogen, and 330 µL of 100% EtOH. Mix by vortexing for 30 s. Incubate the mixture at ~80°C for 16 h.

c. Centrifuge the mixture to pellet the DNA (21,000 × g; 30 min; 4°C) and remove supernatant.

d. Wash pellet in 1 mL of 80% EtOH kept at 4°C for 5 min and remove supernatant.

e. Centrifuge (21,000 × g ;5 min; 4°C) and remove supernatant.

f. Air dry pellets for 5 min at 25°C.

g. Add 20 µL nuclease-free H2O. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the extension products.

11. Gel purification of primer extension products
Next, we perform gel purification of primer extension products to remove excess primer followed by ethanol precipitation.

a. Add 20 µL of 2X loading dye (95% deionized formamide, 18 mM EDTA, 0.02% SDS, xylene cyanol, bromophenol blue).

b. Load 40 µL of the sample in one well of 10% 7M urea slab gel (equilibrated and run in 1X TBE) alongside Low Range ssRNA Ladder. Run gel at 80 V for ~25 min to allow the sample to enter the gel, then increase the voltage to 200 V and run until the bromophenol blue dye front reaches approximately one half of the total length of the gel (~4 cm on an 8 cm gel).

c. Visualize nucleic acids by using SYBR Gold nucleic acid gel stain followed by UV transillumination (Figure 4A).

Note: The amounts of templates of particular library can vary and in some cases the products are not seen on stained gel until the amplification step.

![Figure 4. Primer extension and 3’-adapter ligation: results](image)
d. Excise ssDNA products ~40–80 nt in size from the gel using a sterile scalpel.

e. Elute nucleic acids from gel by crushing gel slices as described in (Vvedenskaya and Nickels, 2020), adding 350 µL of 0.3 M NaCl in 1 X TE Buffer (10 mM Tris-HCl pH 8, 0.1 mM EDTA) to the crushed gel, and incubating the slurry for 10 min at 70°C.

f. Transfer slurry to a Spin-X centrifuge tube filter, centrifuge (14,000 x g; 1 min; 25°C), and transfer eluate into a 1.7 mL tube.

g. Remove the crushed gel from the Spin-X tube and perform steps 11.e-f one more time. Combine eluates (total volume ~700 µL).

h. Add 70 µL of 3 M NaOAc, 7.7 µL of glycogen (10 mg/mL) and 650 mL of isopropanol, mix by vortexing for 30 s, and incubate at –80°C for 16 h.

i. Centrifuge (21,000 x g; 30 min; 4°C) to pellet the DNA and remove supernatant.

j. Wash DNA pellet in 1 mL of 80% EtOH kept at 4°C for 16 h. Centrifuge (21,000 x g; 5 min; 4°C) and remove supernatant.

k. Perform step 11.j two more times.

l. Air dry pellet for 5 min at 25°C.

m. Add 5 µL nuclease-free H2O. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.

Note: Our protocol is aimed at preparation of libraries suitable for high throughput sequencing on Illumina sequencing platform. The ssDNA size selection at step 11.d provides template size range that is suitable for 75-cycle sequencing run.

Note: Gels for distinct primer extension reactions should be stained in separate containers to avoid cross contamination of samples.

Pause point: DNA can be stored at –80°C for ~6 months.

### 3’-Adapter ligation, amplification, and gel purification of libraries

@ Timing: ~18 h for step 12

@ Timing: ~19 h for step 13

@ Timing: ~3 h for step 14

@ Timing: ~20 h for step 15

@ Timing: ~19 h for step 16

The next step of the protocol involves ligation of an oligonucleotide (the “3’ adapter”) to 3’ end of primer extension products prepared in the previous section. After ligation of the 3’ adapter, emulsion PCR is performed, reaction products are separated by gel electrophoresis, and amplicons of the desired lengths are purified by gel extraction. The gel-purified products are then sequenced using an Illumina NextSeq platform.

12. DNA adapter ligation to the 3’ end of the extension products isolated from gel

Note: We have found that use of a two-step ligation procedure improves yields of adapter-ligated products. The first step uses 5’-AppDNA/RNA Ligase, which can join 3’-OH end of DNA substrate and 5’ end of adenylated DNA adapter (AppDNA). The second step uses T4 RNA Ligase 1, which can join 3’-OH end of DNA substrate and 5’ end of DNA adapter that has 5’-phosphoryl group (5’P-DNA). We speculate that the 5’P-DNA, the original substrate
in the adenylation reaction by the Mth enzyme (NEB) is a potential co-reagent in inefficient adapter adenylation reaction.

a. Prepare the first ligation reaction as follows:

| Reagent                                      | Final concentration | Amount |
|----------------------------------------------|---------------------|--------|
| Extension products from step 11.m           | n/a                 | 5 μL   |
| App s1248 (16.9 μM)                          | 0.85 μM             | 0.5 μL |
| 10 X NEB Buffer 1                            | 1 X                 | 1 μL   |
| 50 mM MnCl2                                  | 5 mM                | 1 μL   |
| 5'-AppDNA/RNA Ligase (20 μM)                 | 1 μM                | 0.5 μL |
| Nuclease-free H2O                            | n/a                 | 2 μL   |
| Total                                        | n/a                 | 10 μL  |

b. Incubate for 1 h at 65°C, followed by 3 min at 90°C, cool to 4°C and incubate for 5 min. Use the entire mixture for the second ligation reaction.

c. Prepare the second ligation reaction as follows:

| Reagent                                      | Final concentration | Amount |
|----------------------------------------------|---------------------|--------|
| First ligation reaction from step 12.b       | n/a                 | 10 μL  |
| 10 X T4 RNA Ligase Reaction Buffer           | 1 X                 | 2.5 μL |
| 50% PEG8000                                  | 12%                 | 6 μL   |
| 100 mM DTT                                   | 10 mM               | 2.5 μL |
| 1 mg/mL BSA                                  | 60 μg/mL            | 1.5 μL |
| T4 RNA Ligase 1 (10 U/μL)                    | 10 U                | 1 μL   |
| Nuclease-free H2O                            | n/a                 | 1.5 μL |
| Total                                        | n/a                 | 25 μL  |

d. Pipette the mixture 10 times, briefly spin and incubate at 16°C for 16 h.

13. Isolation of adapter-ligated products by gel size selection

Next, adapter-ligated products are separated from excess adapter using gel size selection and purified by ethanol precipitation.

a. Add 20 μL of 2X loading dye (95% deionized formamide, 18 mM EDTA, 0.02% SDS, xylene cyanol, bromophenol blue).

b. Load the sample in one well on a 10% 7M urea slab gel (equilibrated and run in 1 X TBE) alongside Low Range ssRNA Ladder. Run gel at 80 V for ~25 min to allow the sample to enter the gel, then increase the voltage to 200 V and run until the bromophenol blue dye front reaches approximately one half of the total length of the gel (~4 cm on an 8 cm gel).

c. Visualize nucleic acids using SYBR Gold nucleic acid gel stain followed by UV transillumination (Figure 4B).

Note: Gels for individual ligation mixture should be stained in separate containers.

Note: For alternative options of adapter ligation see: Troubleshooting, Problem 2.

d. Using a sterile scalpel excise 3'-adapter-ligated products. These products will be longer than the primer extension products by 37 nt (~75–120 nt).

Note: The yields of products in the desired size range may be too low to directly visualize on the gel at step 13.c (Figure 4B) for several reasons, including low template yields and co-migration of the templates with the bromophenol blue dye in the loading buffer (see also step 11.c, section “Primer extension reactions and purification of primer extension products”).
e. Elute nucleic acids from gel by crushing gel slices as described in (Vvedenskaya and Nickels, 2020), adding 350 μL of 0.3 M NaCl in 1X TE Buffer (10 mM Tris-HCl pH 8, 0.1 mM EDTA) to the crushed gel, and incubate the slurry for 10 min at 70°C.

f. Transfer the slurry to a Spin-X centrifuge tube filter, centrifuge (14,000 g; 1 min; 25°C) and transfer the eluate into a 1.7 mL tube.

g. Remove the crushed gel from the Spin-X tube and perform steps 13.e-f one more time. Combine eluates (total volume ~700 μL).

h. Add 70 μL of 3 M NaOAc, 7.7 μL of glycogen (10 mg/mL) and 650 μL of isopropanol, mix by vortexing for 30 s, and incubate at −80°C for 16 h.

i. Centrifuge the mixture to pellet DNA (21,000 × g; 30 min; 4°C) and remove supernatant.

j. Wash pellet in 1 mL of 80% EtOH kept at 4°C. Centrifuge (21,000 × g; 5 min; 4°C) and remove supernatant.

k. Perform steps 13.j two more times.

l. Air dry pellets for 5 min at 25°C.

m. Add 15 μL nuclease-free H₂O to DNA pellet. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.

**Pause point**: Purified adapter-ligated products can be stored at −80°C for ~6 months.

14. Control PCR amplification of adapter-ligated products

   Next, a control PCR amplification is performed to estimate the yield and quality of a library generated from the adapter-ligated products.

   a. Prepare control PCR reaction mix as indicated below:

   | Reagent                                   | Final concentration | Amount |
   |-------------------------------------------|---------------------|--------|
   | Adapter-ligated products from step 13.m   | n/a                 | 2 μL   |
   | 5X HF Phusion Buffer with MgCl₂           | 1X                  | 2 μL   |
   | 10 mM dNTP mix                            | 0.2 mM              | 0.2 μL |
   | 2.5 μM Illumina RP1 Primer                | 0.25 μM             | 1 μL   |
   | 2.5 μM Illumina Index Primer (RP11-48)    | 0.25 μM             | 1 μL   |
   | HF Phusion DNA Polymerase, 2 U/μL         | 0.2 U               | 0.1 μL |
   | Nuclease-free H₂O                         | n/a                 | 3.7 μL |
   | Total                                     | n/a                 | 10 μL  |

   b. Perform reactions using cycling conditions below:

   | Steps            | Temperature | Time | Cycles |
   |------------------|-------------|------|--------|
   | Initial Denaturation | 98°C      | 30 s | 1      |
   | Denaturation      | 98°C      | 10 s | 12 cycles |
   | Annealing         | 62°C      | 20 s |        |
   | Extension         | 72°C      | 10 s |        |
   | Final extension   | 72°C      | 5 min  | 1     |
   | Hold             | 4°C      | forever |    |

   c. Mix 10 μL of amplified DNA with 3 μL of 6× Orange DNA load dye and run a non-denaturing 10% TBE gel at the conditions described at step 13.b. Load O’Gene Ruler Ultra Low Range DNA ladder as size standard. Run gel until yellow dye reaches gel bottom.

d. Visualize nucleic acids using SYBR Gold nucleic acid gel stain followed by UV transillumination. Verify library quality by size range and band(s) intensity.

**Note**: Library quality verification can also be performed using an automated electrophoresis system.
15. Amplification of library using emulsion PCR

Next, the library is amplified and purified by use of Micellula DNA Emulsion and Purification Kit (emulsion PCR or ePCR) followed by ethanol precipitation.

**Note:** Use of ePCR reduces the generation of amplicons derived from template switching (Odelberg et al., 1995).

a. Use Micellula DNA Emulsion and Purification Kit to prepare ePCR reaction using manufacturer recommendations.

| Reagent                                      | Final concentration | Volume |
|----------------------------------------------|---------------------|--------|
| Adapter-ligated products from step 13.m      | n/a                 | 1–2 μL |
| 5 X Detergent-Free HF Phusion Buffer with MgCl2 | 1 X                 | 10 μL  |
| 0.1 mg/mL BSA                                | 5 μg/mL             | 2.5 μL |
| 10 mM dNTP mix                               | 400 μM              | 2 μL   |
| 10 μM Illuma RP1 Primer                      | 0.5 μM              | 2.5 μL |
| 10 μM Illuma Index Primer (RP11-48)          | 0.5 μM              | 2.5 μL |
| HF Phusion DNA Polymerase, 2 U/μL            | 0.04 U              | 1 μL   |
| Nuclease-free H2O                            | n/a                 | 27.5–28.5 μL |
| **Total**                                    | n/a                 | 50 μL  |

b. Perform ePCR reactions using the following thermocycler conditions:

| Steps             | Temperature | Time | Cycles |
|-------------------|-------------|------|--------|
| Initial Denaturation | 95°C       | 10 s | 1      |
| Denaturation       | 95°C       | 5 s  | 20 cycles |
| Annealing          | 60°C       | 5 s  |        |
| Extension          | 72°C       | 15 s |        |
| Final extension    | 72°C       | 5 min| 1      |
| Hold               | 4°C        | forever| |

c. Recover ePCR amplicons by breaking the emulsion and purifying DNA according to the manufacturer’s recommendations. Elute DNA from the filter with 150 μL of elution buffer, add 15 μL 3 M NaOAc, 1.5 μL glycerol (10 mg/mL) and 500 μL 100% EtOH.

d. Precipitate the eluate at −80°C for 16 h.

e. Centrifuge the mixture to pellet DNA (21,000 × g; 30 min; 4°C) and remove the supernatant.

f. Wash pellet in 1 mL of 80% EtOH kept at 4°C. Centrifuge (21,000 × g; 5 min; 4°C) and remove supernatant.

g. Perform step 15.f two more times.

h. Air dry pellets for 10 min at 25°C.

i. Add 20 μL nuclease-free H2O to DNA pellet. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.

d. Precipitate the eluate at −80°C for 16 h.

e. Centrifuge the mixture to pellet DNA (21,000 × g; 30 min; 4°C) and remove the supernatant.

f. Wash pellet in 1 mL of 80% EtOH kept at 4°C. Centrifuge (21,000 × g; 5 min; 4°C) and remove supernatant.

g. Perform step 15.f two more times.

h. Air dry pellets for 10 min at 25°C.

i. Add 20 μL nuclease-free H2O to DNA pellet. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.

d. Precipitate the eluate at −80°C for 16 h.

e. Centrifuge the mixture to pellet DNA (21,000 × g; 30 min; 4°C) and remove the supernatant.

f. Wash pellet in 1 mL of 80% EtOH kept at 4°C. Centrifuge (21,000 × g; 5 min; 4°C) and remove supernatant.

g. Perform step 15.f two more times.

h. Air dry pellets for 10 min at 25°C.

i. Add 20 μL nuclease-free H2O to DNA pellet. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.

d. Precipitate the eluate at −80°C for 16 h.

e. Centrifuge the mixture to pellet DNA (21,000 × g; 30 min; 4°C) and remove the supernatant.

f. Wash pellet in 1 mL of 80% EtOH kept at 4°C. Centrifuge (21,000 × g; 5 min; 4°C) and remove supernatant.

g. Perform step 15.f two more times.

h. Air dry pellets for 10 min at 25°C.

i. Add 20 μL nuclease-free H2O to DNA pellet. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.
e. Elute nucleic acids from gel by crushing gel slices as described in (Vvedenskaya and Nickels, 2020), adding 350 mL of 0.3 M NaCl in 1 X TE Buffer (10 mM Tris-HCl pH 8, 0.1 mM EDTA) to the crushed gel, and incubating the slurry for 2 h at 37°C.

f. Transfer the slurry to a Spin-X centrifuge tube filter, centrifuge (14,000 × g; 1 min; 25°C) and transfer the eluate into a 1.7 mL tube.

g. Remove the crushed gel from the Spin-X centrifuge tube filter and repeat steps 16.e-f.

h. Combine eluates from steps 16.f-g (total volume ~700 mL).

i. Add 70 mL of 3 M NaOAc, 7.7 mL of glycogen (10 mg/mL) and 650 mL of isopropanol, mix by vortexing for 30 s, and incubate at ~80°C for 16 h.

j. Centrifuge the mixture to pellet DNA (21,000 × g; 30 min; 4°C) and remove supernatant.

k. Wash DNA pellet 3 times in 1 mL of 80% EtOH kept at 4°C. Centrifuge (21,000 × g; 5 min; 4°C) and remove supernatant after each wash step.

l. Air dry pellets for 10 min at 25°C.

m. Add 20 mL nuclease-free H2O. Incubate at 25°C for 5 min and pipette the sample 10–15 times to ensure the full resuspension of the nucleic acids.

n. Measure the concentration of the libraries using Qubit dsDNA HS Assay kit prior to sequencing. Run control 10% TBE gel to check the quality of final library. To ensure accurate quantification of each library prior to sequencing we recommend use of an automated electrophoresis system. We routinely use an Agilent TapeStation system to assess the quality of our libraries (Figure 6).

o. Sequence library. See (Winkelman et al., 2020) for a detailed description of the analysis of the sequencing data.

**Note:** Any automated electrophoresis system, Bioanalyzer Systems, Fragment Analyzer Systems or TapeStation Systems, can be used for efficient and accurate analysis of libraries prepared for high throughput sequencing.

**Note:** For additional details on how to improve the yield and the quality of the final library see: Troubleshooting, Problems 3–5.

**Note:** Barcoded libraries generated by XACT-seq are suitable for sequencing on Illumina NextSeq platform in high-output mode. Illumina PCR forward and index primers from TruSeq Small RNA Sample Prep Kits are used. Sequencing primers provided by Illumina contain a mixture of several oligos. To avoid potential complications due to the presence of a mixture...

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**Figure 5.** ePCR results: analysis by standard polyacrylamide gel electrophoresis (PAGE)

PAGE analysis of amplicons generated in ePCR reactions before (step 15.i) and after gel purification (step 16.n, section “3’-Adapter ligation, amplification, and gel purification of libraries”). DNA products of 160–180 bp in size are the amplicons of interest. DNA band of ~130 bp in size are amplicons of unknown origin. M, O’Gene Ruler Ultra Low Range DNA ladder.
of primer sequences in the Illumina sequencing reagents we use a custom sequencing primer, s1115.

EXPECTED OUTCOMES

Representative results of primer extension and 3′-adapter ligation reactions are shown in Figure 4. Representative results of ePCR reactions are shown in Figures 5 and 6.

LIMITATIONS

The protocol provided above describes use of XACT-seq to analyze transcription complexes in the context of plasmid-borne sequences in E. coli. Application of XACT-seq to analyze transcription complexes in the context of the bacterial chromosome would require modifications to the protocol (e.g., inclusion of a DNA fragmentation step prior to IMAC, modification of the adapter ligation steps, and modification of the ePCR step).

The transcription complex can sample pre-translocated, post-translocated, reverse-translocated, and hyper-translocated states (Belogurov and Artsimovitch, 2019; Larson et al., 2011). XACT-seq, when used on its own, is unable to define the translocation state of a transcription complex. This limitation of XACT-seq can be overcome by combining XACT-seq and NET-seq for parallel analysis of RNAP active-center A-site positions and RNA 3′ end positions, respectively, each with single-nucleotide resolution.

We used the model bacterium E. coli to develop and first apply XACT-seq, but XACT-seq could be applied to organisms with more complex genomes. Bpa has previously been incorporated into RNAP subunits in yeast (Chen et al., 2007) and can be incorporated into proteins in mammalian cells (Hino et al., 2005). Methods for UV-irradiation, DNA fragmentation, RNAP purification, and sequencing library preparation would be similar for each organism. Application of XACT-seq to other organisms requires development and validation of a Bpa-containing RNAP derivative that, upon UV-irradiation, crosslinks to DNA at a precise position relative to the RNAP active center in a manner analogous to the reagent used in this work (E. coli RNAP, β(R1148Bpa).

TROUBLESHOOTING

Problem 1

Number of individual transformants obtained prior to steps 1.e and 3.d in section “Introduction of plasmids into E. coli cells” is low.
Potential solution
Check the quality of plasmid DNA used in transformation and perform additional purification to remove contaminating salts/organics. Optimize the concentration of plasmid DNA used in the transformation. Use larger volume of competent cells for transformation. Perform additional or multiple transformations using the same stock of plasmid and combine transformants. Repeat transformation with freshly generated competent cells.

Problem 2
Low yield of ePCR products (steps 15, section “3’-Adapter ligation, amplification, and gel purification of libraries”) because of low adapter ligation efficiency.

Potential solution
Other ligases can be tested in steps 12.a-d in section “3’-Adapter ligation, amplification, and gel purification of libraries.” The CircLigase ssDNA ligase (Lucigen) joins 3’-OH end of DNA substrate and 5’ end of DNA adapter that has 5’-phosphoryl group (5’P-DNA). The T4 DNA ligase HC (Thermo Fisher Scientific) joins 3’-OH end of DNA substrate and 5’-P end of double stranded adapter that has overhang of a randomized sequence to provide an efficient annealing to the substrate. Each ligase requires an optimization of reaction conditions.

Problem 3
Low yield of ePCR products (step 15, section “3’-Adapter ligation, amplification, and gel purification of libraries”) because of low input DNA in ePCR reaction.

Potential solution
Increase amounts of adapter-ligated products used in the ePCR reaction. Perform multiple ePCR reactions. Increase the number of amplification cycles to 30.

Problem 4
Presence of non-specific products in the final library (step 16.n, section “3’-Adapter ligation, amplification, and gel purification of libraries”, Figures 5 and 6).

Potential solution
Perform a second gel extraction step of amplified templates as described in step 16, section “3’-Adapter ligation, amplification, and gel purification of libraries”.

Problem 5
Presence of non-specific ~130-bp amplicon in the final library (step 16.d, section “3’-Adapter ligation, amplification, and gel purification of libraries”, Figures 5 and 6).

Potential solution
Carry over of excess primer extension oligo (step 11, section “Primer extension reactions and purification of primer extension products”) or excess 3’ adapter oligo (step 13, section “3’-Adapter ligation, amplification, and gel purification of libraries”) as a result of gel staining at high agitation or prolonged staining time can result in non-specific ~130-bp amplicon in the final library. Hence, care should be taken to avoid an oligo carry over at library gel selection step. One of the potential solutions is loading an oligo on gel in a well separated from the library by one empty well. Run the gel as described in the sections above and separate the part containing an oligo. Stain with SYBER GOLD staining solution and define the position of oligo. Use this as a guide to separate part of the gel containing the library from the part of the gel containing excess oligo. Stain two gel pieces with SYBER GOLD staining solution in separate containers. Use a gel part containing the library for elution of library templates, and a gel part containing excess oligo to control that no traces of oligo left on the top piece. The procedure requires optimization but overall is very efficient and at standardized conditions can be done only one time. Also, suboptimal PCR reaction conditions can result
in non-specific products. To avoid this, optimize PCR reaction components and amplification conditions.

**RESOURCE AVAILABILITY**

**Lead contact**

Additional information and requests for resources and reagents should be directed to and will be fulfilled by the lead contact, Bryce Nickels (bnickels@waksman.rutgers.edu).

**Materials availability**

Plasmids and strains are available upon request.

**Data and code availability**

Unprocessed sequencing reads have been deposited in the NIH/NCBI Sequence Read Archive under the study accession number PRJNA615362. Source code and documentation for analysis of sequencing data are provided at https://github.com/NickelsLabRutgers/XACT-seq.

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**AUTHOR CONTRIBUTIONS**

C.P. designed research, performed research, and wrote the manuscript. I.O.V. designed research, performed research, and wrote the manuscript. B.E.N designed research, acquired funding, supervised the project, and wrote the manuscript.

**DECLARATION OF INTERESTS**

The authors declare no competing interests.

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