Small RNA analysis using the Genome Sequencer™ FLX system

The Genome Sequencer™ FLX System from 454 Life Sciences™ and Roche Applied Science is a versatile sequencing platform suitable for a wide range of applications, including de novo sequencing and assembly of genomic DNA, transcriptome sequencing, metagenomics analysis and amplicon sequencing. The Genome Sequencer FLX is built upon 454 Sequencing™ technology. One application in which the technology has had a considerable impact is small RNA analysis.

Small RNAs, including microRNAs, endogenous siRNAs and Piwi-interacting RNAs, are a diverse collection of molecules with several important biological functions. Genome Sequencer technology is ideally suited for the analysis of small RNA, and several small-RNA studies that use this technology have been published1–8. With the Genome Sequencer FLX System (Fig. 1), each small RNA molecule within a mixture is sequenced individually. The ability to quickly sequence tens to hundreds of thousands of individual small RNA molecules without cloning into bacteria allows quick identification and quantification of the small RNAs in a sample.

Preparing small RNA for sequencing

Preparation of samples for sequencing begins with the generation of double-stranded cDNA from the small RNA. To facilitate the study of small RNAs, these molecules are typically modified with flanking adaptors and converted to cDNA9 using standard protocols. The resulting cDNA, which is suitable for analysis, comprises the small RNAs, typically 20–30 bp in length, flanked by common adaptors of ~20 bp. An online protocol for the preparation of the double-stranded cDNA is available from the David Bartel lab at the Whitehead Institute (http://web.wi.mit.edu/bartel/pub/protocols_reagents.htm) and from the Victor Ambros lab at Dartmouth Medical School (http://banjo.dartmouth.edu/lab/MicroRNAs/Ambros_microRNAcloning.htm).

There are two general methods for preparing these double-stranded cDNAs for sequencing with the Genome Sequencer FLX System. In one method, the double-stranded cDNA is used as starting material for the standard Genome Sequencer FLX DNA library preparation protocol. The Genome Sequencer process–specific A and B adaptors (44-mer adaptors) are blunt-end ligated onto the double-stranded cDNA copy of the small RNA molecules. The resulting double-stranded library is transferred to the emulsion PCR (emPCR™) step for clonal amplification and subsequent sequencing of the individual small RNA molecules (Fig. 2). Sequence data derived from this method have been presented in several publications5–8.

In the second method, Genome Sequencer process–specific A and B adaptors (Fig. 3) are appended onto the double-stranded cDNA copy of the small RNA during a PCR step. In this method, the PCR amplification primers are a fusion primer pair comprising the ~20-bp small RNA adaptor and a 19-bp fixed sequence (Primer A or Primer B) that

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serve as priming sites during the clonal amplification and the sequencing reaction. The double-stranded library of small RNA molecules is clonally amplified via emPCR and sequenced. Several studies have used this method \(^2,4,7\), and detailed protocols are available in these publications.

**Preparation of single-stranded tailed templates for sequencing**

A variant of the method for appending Genome Sequencer process–specific A and B adaptors onto small RNA by PCR (the second method described above) was developed by Wendy Johnston in the David Bartel lab. This protocol generates single-stranded 19-mer tailed templates that can be used directly in emPCR and sequencing. The protocol, presented below, is a modification of a published method \(^10\). Sequence data derived by this method have been presented in several publications \(^3,7\).

**Protocol for preparing templates**

Prepare PCR templates with one tailed primer and one poly(A)-extend ed tailed primer having the following sequences: 5’ primer, 5’-GCCTCCTCGGCGCCATCAG-3’; 3’ primer, 5’-AAAAAAAAAAAAAAAAAAAATSp18/GCCTTGCCCA GCCCGCTCAGTATTGATGGTGCCTACAG-3’.

- Prepare a 100-µl reaction as follows: 0.5 µl of template
  
  0.75 µl of 100 µM 5’ primer
  
  0.75 µl of 100 µM 3’ primer
  
  1 µl of 7aq polymerase (5 units/µl)
  
  10 µl of 10× dNTPs (1×: 0.2 mM each dNTP)
  
  10 µl of 10× PCR buffer (1×: 10 mM Tris-HCl (pH 8.4), 50 mM KCl, 1.5 mM MgCl\(_2\), 0.01% gelatin)
  
  77 µl H\(_2\)O

- Run 12–16 cycles of PCR (each cycle: 1 min 94 ºC, 1 min 50 ºC, 1 min 72 ºC), according to how quickly the band appears. Use the lowest possible number of cycles.

- Stop PCR with the addition of EDTA to a final concentration of 2 mM.

- Precipitate the DNA: add NaCl to a final concentration of 300 mM, add 2 volumes of ethanol and place reaction mix at –20 ºC for at least 2 h.

Prepare a 90% formamide, 8% acrylamide gel (1.5 mm thick); use 7.5 inch × 7.5 inch plates and 20-well comb (each lane is 0.5 cm wide).

- In a beaker, combine 10 ml 10× Tris-borate-EDTA buffer (TBE), 400 mg bisacrylamide, 90 ml formamide and 7.6 g acrylamide. Mix using stir plate until dissolved.

- Filter-sterilize, using suction, to remove any particles and oxygen. Store in a light-blocking bottle.

- For a 50-ml gel, use 500 µl of 10% ammonium persulfate and 150 µl of TEMED. Mix vigorously. Polymerization is very slow; it is best to allow the gel to polymerize overnight.

- Prepare 1× formamide loading dye: 900 µl of formamide, 100 µl of 1× TBE, 10 µl of bromophenol blue and xylene cyanol dye solution.

Gel-purify PCR products.

- Centrifuge precipitations for 30 min at full speed.

- Resuspend in 12–15 µl of 1× formamide loading dye.

- For each marker lane, add 10 µl of 1× formamide loading dye to 2 µl of 1 µg/µl 10-bp ladder in water.

- Pre-run gel at 20 watts for 45 min.

- Heat samples, including ladder, for 10 min at 85 ºC.

- Load samples on gel, with a ladder next to the sample. Run gel at 20 watts until bromophenol blue runs off the gel.

- Place the gel on plastic wrap in a container. Stain with a solution of 1× TBE–ethidium bromide (~20 µl ethidium bromide and 150 µl of TEMED). Mix vigorously. Polymerization is very slow; it is best to allow the gel to polymerize overnight.

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~ 45 min without shaking (with shaking, ~25 min). (If concerned about cross-contamination, load each sample on a different gel. Otherwise, load multiple samples on the same gel, each with a marker lane, then cut the gel into pieces and stain each piece separately.)

• Visualize staining with UV light. Two bands should be present: one at 130 bp and one at 100 bp. Cut the gel between the two bands (at about 115 bp), approximately 15–20 bp above the 130-bp band, and approximately 15–20 bp below the 100-bp band.

• Elute the two pieces separately in 450 µl of 300 mM NaCl overnight.

• Ethanol-precipitate higher bands as for the DNA template above. Wash pellet in cold 70% ethanol; centrifuge again. Evaporate any remaining formamide (30–60 min) by vacuum concentration. Resuspend pellet in 20 µl of water.

The sample is ready for sequencing.

Summary
Sequencing with the Genome Sequencer FLX System provides a quick, efficient and cost-effective means of sequencing small RNAs. Preparing small RNAs for sequencing is straightforward: the starting point for sample preparation is the standard small RNA–to–cDNA protocol. With multiple sequencing formats available, you can tailor the number of clonal small RNA reads per sample to experimental goals. And, unlike other next-generation sequencing technologies, the read lengths in the system are long enough to allow a complete read-through of the standard cloning adaptors in addition to the small RNA, thereby providing a quality assurance of your sequence results.

Additional information about the Genome Sequencer System is available from Roche Applied Science (http://www.genome-sequencing.com). Genome Sequencer, 454, 454 Life Sciences, 454 Sequencing, and emPCR are trademarks of 454 Life Sciences Corporation, Branford, Connecticut, USA. License disclaimer information is available online (http://www.genome-sequencing.com).


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