3' End Sequencing Library Preparation with A-seq2

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Abstract

Studies in the last decade have revealed a complex and dynamic variety of pre-mRNA cleavage and polyadenylation reactions. mRNAs with long 3' untranslated regions (UTRs) are generated in differentiated cells whereas proliferating cells preferentially express transcripts with short 3'UTRs. We describe the A-seq protocol, now at its second version, which was developed to map polyadenylation sites genome-wide and study the regulation of pre-mRNA 3' end processing. Also this current protocol takes advantage of the polyadenylate (poly(A)) tails that are added during the biogenesis of most mammalian mRNAs to enrich for fully processed mRNAs. A DNA adaptor with deoxyuracil at its fourth position allows the precise processing of mRNA 3' end fragments for sequencing. Not including the cell culture and the overnight ligations, the protocol requires about 8 h hands-on time. Along with it, an easy-to-use software package for the analysis of the derived sequencing data is provided. A-seq2 and the associated analysis software provide an efficient and reliable solution to the mapping of pre-mRNA 3' ends in a wide range of conditions, from 10⁶ or fewer cells.

Video Link

The video component of this article can be found at https://www.jove.com/video/56129/

Introduction

The capture and sequencing of mRNA 3' ends allows the study of mRNA processing and the quantification of gene expression. Due to their poly(A) tails, eukaryotic mRNAs can be efficiently purified from total cell lysates with bead-immobilized oligo-deoxythymidine (oligo(dT)) molecules, which can also prime cDNA synthesis. However, this approach has two drawbacks. First, stretches of A’s that are internal to transcripts can also prime cDNA synthesis, resulting in spurious poly(A) sites. Second, homogeneous poly(A) stretches pose specific challenges for sequencing, aside from not being informative for transcript identification. Various approaches have been proposed to circumvent these limitations, such as reverse transcription through poly(A) tails followed by RNase H digestion (3P-seq⁻¹), use of a custom sequencing primer ending in 20 Ts (2p-seq⁻²), preselection of RNA fragments with poly(A) tails of over 50 nucleotides with a CU₅₅ primer followed by RNase H digestion (3'READS⁻³), and the use of an oligo-dT primer that contains the 3' adapter in a hairpin (A-seq⁻⁴).

The recently developed A-seq2 method⁵ aims to bypass sequencing through poly(A) and at the same time to minimize the proportion of dimers that are generated by self-ligation of adapters, particularly occurring when the molar concentration of adapters outweighs the insert concentration. This problem can be eliminated when both adapters are ligated to the same type of polynucleotide ends as in A-seq2, where the 3' adapters are ligated to the 5' end of RNA fragments and the 5' adapters to 5' ends of the cDNAs after reverse transcription. The method is more convenient than our previously proposed A-seq - in which sequencing was in the 5'-to-3' direction thereby requiring precisely controlled RNA fragmentation-, while maintaining a high accuracy of poly(A) site identification. Around 80% of the sequenced reads in typical samples map uniquely to the genome and lead to identification of over 20,000 poly(A) site clusters, more than 70% of which overlapping with annotated 3'UTRs.

In brief, the A-seq2 protocol starts with mRNA fragmentation and ligation of reverse-complement 3' adapters to the 5' ends of RNA fragments. Poly(A)-containing RNAs are then reverse transcribed with a 25 nucleotide (nt) long oligo(dT) primer that contains an anchor nucleotide at the 3' end, a dU at position 4 and a biotin at the 5' end, allowing binding of the cDNA to magnetic streptavidin beads. Most of the primer, including the biotin, is removed from the cDNA by cleavage at dU by the USER enzyme mix, containing Uracil DNA glycosylase (UDG) and the DNA glycosylase-lyase Endonuclease VIII. This reaction leaves intact ends for the ligation of a 5' adapter, and the three Ts left after cleavage remain to mark the location of the poly(A) tail. Because both 5' and 3' adapters are attached by ligation to recipient 5' ends, no adapter dimers are generated. Four nucleotide random-mers introduced at the beginning of reads allows cluster resolution on state-of-the-art sequencing instruments and can also serve as unique molecular identifier (UMI) for the detection and removal of PCR amplification artifacts. The size of the UMI can be further increased as done in other studies⁶. The protocol generates reads that are reverse complementary to mRNA 3' ends, all starting with a randomized tetramer followed by 3 Ts. Processing of reads that have the 3 diagnostic Ts at their 5' end starts with the correction of PCR amplification artifacts by exploiting the UMIs, removal of 3' adapter sequences, and reverse complementation. Reads that may have
1. Cell Growth and mRNA Isolation

1. Grow cells according to your experimental design in 6-well plates to ~1 x 10^6 cells per well at 80% confluence.
2. Remove the growth medium and wash the cells once with phosphate buffered saline. Directly lyse the cells on the plate by adding 1 mL of lysis buffer from the mRNA-isolation kit. Transfer the viscous lysate into a 15 mL plastic tube with a 1 mL pipette tip. Use a rubber spatula to completely detach the cell material from the plate surface.
3. Shear the lysate containing viscous DNA with a 1 mL syringe attached to a 23 G hypodermic needle by several vigorous up and down movements of the plunger until the lysate is no longer viscous. Point the syringe needle to the center of the bottom to avoid ejecting the lysate out of the tube.
4. Transfer the lysate into a 1.5 mL tube using the syringe. Spin 5 min at 20,000 x g and 4 °C to remove the debris. Use DNA low bind 1.5 mL vials throughout the protocol.
5. While the centrifuge is running, wash 300 µL of resuspended oligo (dT)25 magnetic beads on a magnetic rack. Resuspend and place tubes on a rotating wheel for 10 min.
6. Place the tubes on a magnetic rack. Remove the clear liquid after 2 min. Add 0.8 mL buffer A from the mRNA-isolation kit. Turn the tube by 180° degrees on the rack, 2-3 times. Repeat this washing step once more with buffer A.
7. Wash the beads 2 times with 0.8 mL of buffer B as described in step 1.6.
8. To elute the bound mRNA from the beads, add 33 µL H2O and resuspend the beads. Heat to 75 °C for 5 min on a heated block. Immediately spin the tubes for 1 s and place them on the magnetic rack. Transfer the supernatant to a new tube. Samples can be stored at -80 °C until further use.
9. Add 66 µL alkaline hydrolysis buffer to the 33 µL mRNA (step 1.8), mix and heat for exactly 5 min at 95 °C on a heating block. Immediately chill tubes on ice.
10. Isolate RNA with an RNA cleanup kit.

   NOTE: Confirm the volume; it should be 100 µL.
   1. Add 350 µL RLT buffer from the kit and 250 µL ethanol. Load onto the column and spin for 30 s at 8,000 x g at room temperature (RT). Wash with 500 µL RPE buffer from the kit. Wash with 500 µL 80% ethanol. Spin for 5 min at 20,000 x g to dry the column. Add 36 µL H2O to column and spin for 1 min at 20,000 x g. Discard the column and save the eluate.

2. 5' end Phosphorylation and DNase Treatment

1. Add 5 µL polynucleotide kinase buffer, 5 µL 10 mM ATP, 1 µL ribonuclease inhibitor, 1 µL DNase and 2 µL polynucleotide kinase to samples and incubate at 37 °C for 30 min. Optionally prepare master reaction mixes throughout the protocol by mixing 1.1 volumes x n (n = number of samples) of each component.
2. Change buffer and remove ATP on a spin-column to prevent poly(A) addition in the next step.
   1. Prespin spin-columns at 735 x g for 1 min. Transfer the columns to new 1.5 mL vials and load the kinase reactions onto the columns. Spin the columns 2 min at 735 x g. Discard the columns and place the tubes with collected reactions on ice or store at -80 °C.

3. Blocking 3' Ends with Cordycepin Triphosphate

NOTE: It is essential to block the 3’ ends of RNA fragments to avoid their concatemerization in the subsequent ligation reactions. 3’ ends that are not already blocked by a (cyclic) phosphate after hydrolysis are treated by addition of a 3’ dATP (cordycepin triphosphate) chain terminator nucleotide with the aid of poly(A) polymerase. Here, yeast poly(A) polymerase (yPAP), that was expressed and purified as described in was used at a concentration of 0.5 mg/mL. Yeast or E. coli PAP both have almost the same activity for addition of 3’dATP and can be purchased commercially (see the Table of Materials).

1. Add 13.5 µL 5x concentrated poly(A) polymerase reaction buffer, 2 µL of 10 mM 3’ dATP, 1 µL RNase inhibitor and 1 µL poly(A) polymerase to the reaction from step 2.2.1. Mix and spin for 1 s. Incubate at 37 °C for 30 min. Add 32.5 µL H2O to each reaction. Purify the RNA as in step 1.10.1. Elute the RNA with 14 µL H2O.
4. Ligation of Reverse 3' Adapters to the 5' End of RNA Fragments

1. Place the reactions in a vacuum concentrator for 10 min to reduce the volume to 6 µL. Add 3 µL 10x T4 RNA ligation buffer, 3 µL 10 mM ATP, 15 µL PEG–8000, 1 µL RNase inhibitor, 1 µL of 0.1 mM reverse complement 3' adapter "revRA3" (see the Table of Materials) and 1 µL high concentration T4 RNA ligase 1, mix.
2. Incubate the reactions at 24 °C for 16 h on a heated mixer with intermittent mixing at 1,000 rpm. Add 70 µL H2O to each reaction and mix. Purify the RNA as in step 1.10.1. Elute the RNA with 14 µL H2O. Samples can be stored at -80 °C at this point.

5. Reverse Transcription (RT)

1. Place the eluates in a vacuum concentrator for 3 min to reduce the volume to 11 µL. Transfer reactions to 200 µL PCR tubes. Add 1 µL 0.05 mM RT primer “Bio-dU-dT25”. Heat for 5 min at 70 °C in a PCR cycler and leave at RT for 5 min.
2. Add 1 µL 10 mM dNTPs, 4 µL 5x reverse transcriptase buffer, 1 µL 0.1 M DTT, 1 µL RNase inhibitor, and 1 µL reverse transcriptase. Mix and heat the reactions for 10 min to 55 °C and 10 min to 80 °C in a PCR cycler. Keep on ice or at -80 °C for longer storage.

6. Digestion with Uracil DNA Glycosylase Enzyme Mix

1. Pipet 100 µL Streptavidin-beads into a 1.5 mL vial, resuspend in 800 µL biotin binding buffer and place on a magnetic rack. Invert tubes 2-3 times. Remove the buffer when clear. Repeat the washing step. Resuspend the beads in 200 µL biotin binding buffer.
2. Add reverse transcription reaction to the beads solution and incubate 20 min at 4 °C on a rotating wheel. Wash the beads 2x with biotin binding buffer. Purify the cleaved cDNA.
3. Add 50 µL H2O, 11 µL of RNase H buffer and 1 µL RNase H to the reactions. Incubate at 37 °C for 20 min. Place tubes on a magnetic rack and transfer the liquid containing the cleaved cDNA to a new tube
4. Purify the cleaved cDNA.
   1. Add 550 µL of buffer PB from the PCR purification kit to the cleavage reactions. Add 10 µL of 3 M sodium acetate, pH 5.2 to lower the pH. Load the reactions on minimal elution spin columns and spin at 17,000 x g for 1 min.
   2. Add 750 µL buffer PE to columns and spin at 17,000 x g for 1 min. Discard the flow-through. Spin the columns at 17,000 x g for 1 min to dry. Transfer the columns to a 1.5 mL vial, add 16 µL H2O and spin at 17,000 x g for 1 min. Place the reactions in a vacuum concentrator for 8 min to concentrate to a volume of 7 µL.

7. Ligation of 5' Adapters to 5' Ends of cDNA

1. To the isolated cDNA, add 3 µL 10x T4 RNA ligase 1 buffer, 3 µL 10 mM ATP, 15 µL PEG–8000, 1 µL 50 µM “revDA5” oligo, and 1 µL high concentration T4 RNA ligase 1. Incubate at 24 °C for 20 h. Add 70 µL H2O to each reaction. Samples can be stored at -20 °C at this point.

8. Pilot PCR, Amplification of Libraries and Size Selection

1. In a pilot reaction, determine the optimal number of PCR cycles to reach library amplification within the exponential phase.
   1. Pipet 25 µL DNA polymerase mix, 20 µL ligation reaction, 2 µL H2O, 1.5 µL 10 µM forward PCR primer (RP1) and 1.5 µL 10 µM reverse PCR index primer into 200 µL PCR tube.
   2. Run the cycler with the following program: 3 min 95 °C, followed by 20 cycles of 20 s 98 °C, 20 s 67 °C and 30 s 72 °C. Collect 7 µL aliquots after 6, 8, 10, 12, 14, 16 and 18 cycles directly from the cycler. Add 1 µL 10x loading buffer (50% glycerol, 0.05% xylene cyanol). Note: Please follow the recommendations of the supplier if using multiplexing when combining barcodes.
3. Separate products in small slots on a 2% agarose gel containing a 1:10,000 dilution of fluorescent green dye. Run the cycler with the following program: 3 min 95 °C, followed by 20 cycles of 20 s 98 °C, 20 s 67 °C and 30 s 72 °C. Collect 7 µL aliquots after 6, 8, 10, 12, 14, 16 and 18 cycles directly from the cycler. Add 1 µL 10x loading buffer (50% glycerol, 0.05% xylene cyanol). Note: Please follow the recommendations of the supplier if using multiplexing when combining barcodes.
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9. Data Processing

NOTE: The resulting sequencing data (in fastq format) are processed with software available in the gitlab repository (https://git.scicore.unibas.ch/zavolan_public/A-seq2-processing). The analysis includes four main steps: (1) downloading the git repository, (2) installation of a virtual environment, (3) setting specific parameters in the configuration file and (4) launching the analysis through ‘snakemake’.

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1. For large-scale PCR reactions, concentrate and desalt the reactions first with a PCR purification kit and separate the products on wide slots on 2% agarose gels in 1x TBE buffer.
2. Use the number of cycles at the beginning of exponential amplification in the pilot reaction for a large-scale PCR reaction with twice the volumes as used for the pilot reaction (Figure 2).
   1. For large-scale PCR reactions, concentrate and desalt the reactions first with a PCR purification kit and separate the products on wide slots on 2% agarose gels in 1x TBE buffer.
   2. Cut out gel slices containing 200-350 nt DNA products. Melt the gel in the chaotropic buffer at RT for up to 30 min. Extract DNA from the gel slices with a gel extraction kit. Do not heat to 50 °C to prevent bias in binding of A-rich DNA.
   3. Submit for sequencing.
   NOTE: Typically, 50 cycles single-read (SR50) are sufficient (see, for e.g., https://www.illumina.com/technology/next-generation-sequencing.html).
repository and a short description is available below. All individual processing steps are accomplished by the execution of publicly available tools, either from external sources or prepared in-house. The computational pipeline depends on an anaconda-based python 3 virtual environment with the snakemake package installed. It runs on machines with Unix-like operating system and was tested in a Linux environment with the CentOS 6.5 operating system installed and 40 GB RAM available. Software dependencies are automatically controlled within the virtual environment. The following publicly available software tools are required and thereby installed together with the environment: snakemake (v3.9.1), fastq toolkit (v0.0.14), STAR (v2.5.2a), cutadapt (v1.12), samtools (v1.3.1), bedtools (v2.26.0).

1. Data pre-processing from reads to cDNAs
   - The sequencing depth may vary between runs and, depending on the instrument, data from one sample can be split over multiple sequence files. If this is the case, concatenate the files that correspond to one sample into a single input file that is used in the following steps.
   1. Convert the file from fastq to fasta format.
   2. Extract reads with a correct structure (3 thymidines at positions 5, 6 and 7 of the read).
      - NOTE: A read that is correctly prepared according to the experimental protocol described above should have the structure (from the 5' end): 4-nucleotide barcode - 3 thymidines - reverse complement of transcript 3' end.
   3. Store the information about the starting tetramer in the description line of the sequence.
      - NOTE: The tetramer serves as a unique molecular identifier (UMI) that facilitates the correction of amplification artifacts later in the analysis.
   4. Remove the first seven nucleotides from the read's 5' end.
   5. Correct for amplification artifacts by keeping only one copy of the reads with the same insert sequence and UMI.
   6. Remove the part of the 3' end that matches the adapter sequence and then reverse complement the sequence. Only proceed with reads that have a minimum length (default: 15 nt).
      - NOTE: Depending on the length of the original mRNA fragment and the number of sequencing cycles, the 3' end of the read may contain part of the 3' adapter, which is removed in this step.
   7. Extract all reads that fulfill the following criteria: maximum 2 unknown nucleotides ('N'), maximum 80% As, and last nucleotide of the read not A. These reads are considered to be of sufficient quality to be used in the analysis.

3. Map the reads to the genome with a tool that handles spliced reads and generates an output file in BAM format.
   1. Create a file with the index of the genome to which the reads should be mapped. For the human genome, this step requires 35 GB of memory (RAM).
   2. Map the reads to the genome.
      - NOTE: (STAR-specific notes) Soft-clipping is disabled in order to force the mapping of the 3' end of each read as this is the nucleotide immediately upstream of the cleavage site.
   3. Remove the part of the 3' end that matches the adapter sequence and then reverse complement the sequence. Only proceed with reads that have a minimum length (default: 15 nt).

4. Convert the BAM to a BED-file. If a read maps to multiple locations, keep only those with the lowest edit distance.
   - NOTE: The copy number of the read mapped at a specific location is used as score. Reads that map to multiple locations are counted fractionally at each location with a weight equal to 1 / number of locations to which a read maps.
   5. Collapse reads that vary by a likely sequencing error. If two distinct reads map to the same location (start and end position of the mappings are identical) and they share the same UMI, consider them as PCR duplicates and keep only one.
   6. Infer all individual pre-mRNA 3' end processing sites.
      - NOTE: An individual read provides evidence for a 3' end when its last four nucleotides are mapped to the genome without error. The position to which the 3' end of the read maps is stored as cleavage site.
   7. Detect 3' end sites that could have originated from internal priming. Define the site as internal priming artifact when the 10 nt downstream of the cleavage site in the genome satisfy one of the following criteria: contains more than six As, contains six consecutive As, or starts with one over million (RPM). Traversing the list from top to bottom, associating lower-ranked sites with higher-ranked sites when their distance to the higher-rank site is not larger than a predefined limit (default: 12 nt). Whenever any of the constituting 3' end site overlaps with an annotated poly(A) signal or has a poly(A) signal directly downstream, the corresponding cluster is marked for further inspection to detect internal priming.
   8. Identify independently regulated poly(A) site clusters.
      - NOTE: The steps described here follow the procedure that was introduced in a prior publication.
      1. Start by collecting individual 3' end processing sites that were obtained in all samples of the study.
      2. Annotate known poly(A) signals in the region of -60 to +10 nucleotides around each individual 3' end processing site.
      3. Identify poly(A) sites expressed above the background in each sample as follows.
         1. Sort the sites by their raw expression within the current sample. Traverse the list of sites from top to bottom, associating lower ranked sites with a higher ranked site if they are located within a predefined distance in the genome (default: 25 nt up- or downstream) from the high-ranking site.
            - NOTE: All low-ranking sites associated with a high-ranking site define a cluster whose expression is the number of reads documenting all of these sites.
         2. Sort these clusters by expression and traverse the list of clusters from highest to lowest expression, determining the expression threshold c at which the percentage of clusters with an annotated poly(A) signal drop below a predefined threshold (default: 90%).
         3. Discard sites from any cluster below the cutoff.
      4. Cluster closely spaced 3' end sites obtained across samples.
         - NOTE: Sort 3' end processing sites first by the number of supporting samples and then by the sum of the normalized read count (reads per million (RPKM)) across samples. Traversing the list from top to bottom, associating lower-ranked sites with higher-ranked sites when their distance to the higher-rank site is not larger than a predefined limit (default: 12 nt). Whenever any of the constituting 3' end site overlaps with an annotated poly(A) signal or has a poly(A) signal directly downstream, the corresponding cluster is marked for further inspection to detect internal priming.
      5. Merge poly(A) site clusters.
         - NOTE: When a cluster is marked as a putative internal priming candidate, it is either merged into a downstream cluster if the two clusters share their poly(A) signals or retained if the most downstream site in the cluster has a poly(A) signal located at a minimum
distance upstream (default: 15 nt). Finally, closely spaced clusters are merged if: (i) they share the same poly(A) signal(s), or (ii) the span of the resulting cluster does not exceed a maximum (default: 25 nt).

6. Store clusters in BED-file format with the total normalized read count from all 3’ end sites in each cluster as score.

**Representative Results**

Poly(A)-containing RNA was isolated from cultured cells, fragmented by alkaline hydrolysis and cDNAs were made by reverse transcription with oligo(dT) primers. The resulting cDNA was immobilized on streptavidin beads, dU was cleaved in the uracil specific excision reaction, adapters were ligated to 5’ and 3’ ends of the cleaved fragment and the inserts were sequenced. Figure 1 depicts a graphical outline of the experiment.

For HeLa and HEK293 cells, $10^6$ cells were sufficient to identify poly(A) sites for the vast majority of protein-coding genes at the end of the procedure. However, for other cell types or tissues it may be necessary to test the saturation in the number of identified poly(A) sites as the number of cells used in the experiment increases. Representative results of the pilot PCR step and of the DNA fragment analysis of the sample before sequencing are shown in Figure 2.

**Figure 3** shows the pre-processing steps of the computational analysis, starting from the fastq file obtained from the sequencer and ending with the quality-checked, adapter-trimmed reads that are ready to be mapped to the genome. **Figure 4** shows the analysis steps that start with the mapping of the reads to the corresponding genome and end with the catalog of mRNA 3’ end processing sites that are identified in a particular sample. When multiple samples are analyzed, additional steps are carried out to match the 3’ end processing sites that were found in individual samples and report their abundance across samples. These steps are shown in **Figure 5**.

Thus, once samples have been sequenced, the analysis of the resulting sequencing read files (in fastq format) through the available processing pipeline is straightforward. After adding the information about the samples to the configuration file, the execution of the pipeline will result in two main types of output files: 1) BED-files with all 3’ end processing sites identified in individual samples (e.g. "sample1.3pSites.noIP.bed.gz"), and 2) a BED-file with all poly(A) site clusters (clusters.merged.bed) across all samples of the study. The output also includes the genome coordinates for all reads from each individual sample (e.g. "sample1.STAR_out/Aligned.sortedByCoord.out.bam") that can later be viewed in a genome browser like IGV. Visual inspection of the read profile(s) generally provides a first glimpse of the distribution of poly(A) sites in the genome and the changes that occur upon the specific perturbations that were carried out in the study. For example, in **Figure 6** the response of a specific gene to the knock-down of the HNRNPC protein is shown.

Summaries of these genome-wide distributions are also provided (Table 1). Specifically, output files in the "counts/annotation_overlap" directory contain fractions of sites that overlap with specific annotated features (from the gtf file provided as input; annotated are: 3’ UTR, terminal exon, exon, intron, intergenic). Finally, for each sample, results of individual processing steps are also saved (e.g. "sample1.summary.tsv"). This includes the numbers of: raw reads in each sample, reads that have the expected structure of the 5’ end, reads that remain after collapsing full PCR duplicates, high-quality reads according to the criteria defined at step 9.2, reads that map uniquely to the genome (after collapsing those that resulted from sequencing errors, see step 9.5), multi-mapping reads (after collapsing those that resulted from sequencing errors, see step 9.5), raw (not clustered) 3’ end processing sites in each sample, raw 3’ end processing sites without potential internal priming candidates, unique 3’ end processing sites from all samples without internal priming candidates, and final set of poly(A) site clusters.
Figure 1: Main steps of the A-seq2 protocol. Individual steps are indicated on the left side of the figure. Insert RNA fragments are depicted as green lines that turn red for cDNA after reverse transcription; adapters are colored in light blue or in orange. Please click here to view a larger version of this figure.

Figure 2: Pilot PCR and final product profile. (a) Aliquots from the PCR reaction were collected at different cycles and separated on 2% agarose gels. Numbers to the left indicate size in nucleotides of the respective bands in the DNA ladder. In this experiment 12 cycles (*) were chosen for the large scale PCR reaction. (b) Example of a sample after size selection run on a fragment size analyzer revealing an average size of around 280 nucleotides. Numbers to the left [FU] indicate relative signal intensity. Please click here to view a larger version of this figure.
Figure 3: Outline of the pre-processing of sequencing reads. The fastq files with reads that are generated by the sequencing instrument-associated software are processed to identify high-quality reads that will be mapped to the corresponding genome. The figure shows the input/output specification of individual steps in the pipeline, with links to the individual steps of the protocol described in section "Data processing". Please click here to view a larger version of this figure.
Figure 4: Outline of sequence read processing, from the step of mapping to the genome to the generation of individual 3' end processing sites. The figure shows the input/output specification of individual steps in the pipeline, with links to the individual steps of the protocol described in section “Data processing”. The main output file that is delivered to the user is marked in bold. Please click here to view a larger version of this figure.
Figure 5: Outline of the steps that are taken to generate clusters of co-regulated 3' end sequencing sites. The figure shows the input/output specification of individual steps in the pipeline, with links to the individual steps of the protocol described in section "Data processing". The main output file is marked in bold. Please click here to view a larger version of this figure.

Figure 6: Example results of the profile of 3' end processing reads along the terminal exon of the NUP214 gene, shown in the IGV genome browser. A-seq2 reads were prepared from two samples of HEK 293 cells, treated either with a control-siRNA or with an HNRNPC siRNA. The reads that documented poly(A) sites that were annotated by the analysis pipeline were saved in BAM format that was used as input to the IGV genome browser. The 3' ends of the read peaks map to mRNA 3' ends that are annotated in Ensembl. The profiles indicate an increased use of the long 3' UTR isoform upon HNRNPC knock-down. Please click here to view a larger version of this figure.
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Disclosures

The authors have nothing to disclose.

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