# Module TSS3:
**CAGE, RAMPAGE, & RNA Pol II X-ChIP-Seq (Advanced)**

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## Lesson Plan:

<table>
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<th>Title</th>
<th>Identifying transcription start sites for peaked promoters using experimental data and conservation</th>
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| Objectives | • Use the placement of the Initiator motif; experimental data from CAGE, RAMPAGE, and RNA Pol II X-ChIP-Seq experiments; and TSS predictions from the Celinker group at modENCODE to identify putative transcription start sites (TSSs)  
• Analyze the multiple sequence alignment (i.e. Drosophila Conservation track) of exon 1 from multiple Drosophila species |
| Pre-requisites | Module TSS1 and Module TSS2 |
| Order | • Explain the experimental techniques used to generate CAGE, RAMPAGE, and RNA Pol II X-ChIP-Seq data, and the Celinker TSS predictions  
• Illustrate how the experimental data and conservation evidence tracks can be used for TSS annotation  
• Investigation 1: Students find the sevenless TSS using CAGE, TSS predictions, RAMPAGE, and RNA Pol II X-ChIP-Seq  
• Investigation 2: Students use the conservation track (ROAST multiple sequence alignments) to compare sequences from multiple Drosophila species  
• Analyze all of the evidence in order to identify the best TSS |
| Class Instruction | • Discuss the questions: What is transcription? How can our understanding of transcription initiation be used to find a TSS?  
• Use the genome browser to introduce TSS evidence tracks  
• Conclude by challenging students to think about these questions:  
  o For each of the evidence tracks you have investigated, how might a broad promoter differ from a peaked promoter?  
  o If a gene has multiple isoforms, does it have to have multiple transcription start sites? |
INTRODUCTION

This module will use the *Drosophila melanogaster* sevenless gene (sev) to illustrate how different experimental techniques (CAGE, RAMPAGE, RNA Pol II X-ChIP-Seq), TSS predictions from the Celniker group at modENCODE, and sequence conservation can be used to identify transcription start sites (TSSs).

INVESTIGATION 1: IDENTIFY THE LOCATION OF TSS USING CAGE, RAMPAGE AND RNA POL II X-CHIP-SEQ DATA

SUMMARY OF CAP ANALYSIS OF GENE EXPRESSION (CAGE)

The relative abundance of mRNA transcripts can be determined by many different methods such as RNA-Seq, quantitative PCR, or Serial Analysis of Gene Expression, which you may have studied in class. However, these methods tend to show a bias toward the body of the transcript. Hence, specialized techniques are needed to isolate sequences associated with the 5' and 3' ends of the transcript.

One technique that can be used to identify a TSS is Cap Analysis of Gene Expression (CAGE). This technique takes advantage of the fact that there is a unique nucleotide at the 5' end of the newly synthesized eukaryotic mRNA, the “5’ cap”. This 5’ cap can be chemically modified, allowing the capped mRNAs to be purified away from RNAs that lack a cap.

The capped mRNAs are first converted to cDNAs by the enzyme reverse transcriptase, then cleaved with an enzyme, leaving short fragments (~27 bp) that correspond to the 5' end of the mRNAs. These short fragments are sequenced, and the resulting sequences (“reads”) are mapped against the genome assembly to identify the TSS. In addition to identifying the TSS, CAGE results can also be used to estimate transcript abundance — that is, relative expression levels.

In Investigation 1, we will first use CAGE data to help identify the TSS. We will also use the TSS predictions produced by the Celniker group at modENCODE to identify the TSS. The Celniker TSS predictions were based on the analysis of experimental data from three methods: CAGE, 5’ Rapid amplification of cDNA ends (5’ RACE), and 5’ Expressed Sequence Tags (ESTs).
EXAMINING THE CAGE DATA FOR THE \textit{SEVENLESS} GENE USING THE UCSC GENOME BROWSER MIRROR

1. Open a new web browser window and go to the Genomics Education Partnership (GEP) UCSC Genome Browser Mirror at \url{http://gander.wustl.edu/} (Figure 1). This genome browser was developed by the Genome Bioinformatics Group at the University of California Santa Cruz (UCSC), and customized by the GEP to facilitate the annotations of multiple Drosophila species.

Figure 1 Access the Genome Browser using the “Genome Browser” link.

2. To navigate to the genomic region surrounding the \textit{sevenless} (\textit{sev}) gene in \textit{Drosophila melanogaster}, select “\textit{D. melanogaster}” under the “genome” field, select “Aug. 2014 (BDGP Release 6 + ISO1 MT/dm6)” under the “assembly” field, and then enter “\textit{sev}” under the “search term” field. Click on the “GO” button (Figure 2).

Figure 2 Change the settings on the Genome Browser Gateway page to navigate to the \textit{sev} gene in the \textit{D. melanogaster} release 6 assembly.

3. This region on the X chromosome, denoted “\texttt{chrX:11,071,441-11,086,299}” contains the entire \textit{sev} gene. The suffix “-RA” corresponds to the name of the isoform that is...
associated with the gene. Hence sev-RA corresponds to the A isoform of the sev gene. This gene only has one isoform.

4. The sev gene is on the minus strand. To make it easier to interpret the evidence tracks, we will reverse complement the entire chromosome sequence. Click on the “reverse” button located in the display controls below the Genome Browser image (Figure 3).

Figure 3  Click on the “reverse” button to reverse complement the entire chrX sequence (red arrow).

5. Because the Genome Browser remembers the previous display settings, we will hide all the evidence tracks and then enable only the subset of tracks that we need. Click on the “hide all” button located below the Genome Browser image. Then, configure the display modes as follows:

- Under “Mapping and Sequencing” Tracks
  - Base Position: **full**
  - Click on the blue “Short Match” link
    - Enter “**TCAKTY**” (i.e. the Initiator motif) into the “Short (2-30 base) sequence” text box
    - Change the “Display mode” to “**pack**”
    - Click on the “**Submit**” button

- Under “Gene and Gene Prediction” Tracks
  - FlyBase Genes: **pack**
• Under “Expression and Regulation”
  o TSS (Celniker) (R5): pack
  o Click on the blue “Combined modENCODE CAGE TSS” link
    • Change the “Maximum display mode” to “full”
    • Scroll down to the “Select views” section, and change the “Peaks” display mode to “pack”
    • In the “List Subtracks” section, uncheck the box for “modENCODE CAGE (Plus)”
    • Scroll up to the top of the page, and then click on the “Submit” button to update the display.

Note: Although we have set the display mode for the “Base Position” track to “full”, we will need to zoom in further to see the nucleotide sequence and the amino acid translations. Enter “chrX:11,086,293-11,086,305” into the “enter position or search terms” text box, and then click on the “go” button.

Note that this exercise requires you to examine the region near exon 1 to determine where transcription begins (Figure 4). The start codon for sev (where translation begins) is in exon 2.

Q1. Based on the “FlyBase Genes” track, what is the coordinate of the transcription start site?

Q2. Based on the Transcription Start Sites (Celniker) (R5) track, what is the coordinate of the transcription start site?

Q3. Is the Celniker TSS prediction consistent with the location of the Inr motif? Why or why not? [Remember that the Initiator (Inr) motif TCAKTY spans the TSS from -2 to +4, and transcription is predicted to start at the A.]
Q4. Based on the Combined modENCODE CAGE TSS track, what is the coordinate of the transcription start site? Remember that this track shows the density of the first base of the CAGE reads, which corresponds to the 5’ end of the mRNA.

Q5. Zoom out 10x, and then zoom out another 3x. Compare the CAGE read density for the TSS that you just identified for the sev gene, and the CAGE read densities for other positions in the region. How many positions within this region have CAGE read densities that are similar to the CAGE read density for the TSS of the sev gene? Do any of these potential TSSs overlap with the Inr motif?
RNA Annotation and Mapping of Promoters for the Analysis of Gene Expression (RAMPAGE)

Additional evidence for the TSS locations can be gathered from RAMPAGE (RNA Annotation and Mapping of Promoters for the Analysis of Gene Expression) data. This technique allows for the identification of the putative TSS with base-pair resolution, and it generally has higher specificity (i.e. signal to noise ratio) than CAGE. This is because whereas CAGE generates short sequences (~27 bp) from capped mRNAs, RAMPAGE allows for sequencing of full-length cDNAs. The sequencing strategy enriches for 5’ sequences by combining two 5’-selection approaches - template-switching and cap-trapping – then generates paired-end reads using Illumina and HiSeq technology. Compared to the short sequences from CAGE, the long paired-end reads from RAMPAGE can be more easily mapped to the genome assembly, thereby facilitating the identification of the TSS.

Importantly, if there are multiple TSSs for a gene (characteristic of genes with a broad promoter), RAMPAGE will quantify how frequently each TSS is used. Batut and colleagues also developed software for analyzing RAMPAGE data. The combination of the experimental and computational techniques make RAMPAGE a powerful method for identifying TSSs.

To examine the RAMPAGE evidence tracks, we will make the following changes to the display modes of the Genome Browser:

- Under “Expression and Regulation”
  - TSS (Celniker) (R5): hide
  - “Combined modENCODE CAGE TSS: hide
  - Click on the blue “Combined RAMPAGE TSS (R5)” link
    - Scroll down to the “Select views” section
      - Change the “Peaks” display mode to “pack”
      - In the “List subtracks” section, uncheck the box for “RAMPAGE (Plus)”
      - Scroll up to the top of the page, and then click on the “Submit” button
  - Click on the blue “RAMPAGE TSS Read Density (R5)” link
    - Change the “Display mode” field to “full”
    - Scroll down to the “Select subtracks by strand and stage” section
• Click on the “-” button at the top left corner to unselect all the datasets (Figure 5)

• Select the “Minus” strand checkboxes for the following developmental stages: Embryos 10hr, Embryos 24hr, L1, L2, Adult Females 5d, Adult Males 5d

• Scroll up to the top of the page, and then click on the “Submit” button to update the Genome Browser display

Figure 5 Configure the “RAMPAGE TSS Read Density (R5)” track to show only the data for six developmental stages on the minus strand. Click on the “-” button at the top left corner to unselect all the evidence tracks (red arrow). Select the checkboxes for the developmental stages of interests underneath the “Minus” column label. (The … denotes additional stages that were omitted from the screenshot.)

• Enter “chrX:11,086,146-11,086,477” into the “enter position or search terms” text box and then click on the “go” button.
Figure 6  The RAMPAGE read density for all samples combined (dark red), and for six developmental stages (bright red) of D. melanogaster.

Q6. How many TSSs are supported by the RAMPAGE evidence track (Figure 6)?

Q7. Do all of the developmental stages shown use the same TSS?

To ascertain whether RAMPAGE identified additional TSS in the other developmental stages, we will reconfigure the “RAMPAGE TSS Read Density (R5)” track to show the minus strand RAMPAGE data for all of the available developmental stages.

- Under “Expression and Regulation”
  - Click on the blue “RAMPAGE TSS Read Density (R5)” link
    - Scroll down to the “Select subtracks by strand and stage” section
    - Click on the “+” button underneath the “Minus” column label (Figure 7)
    - Scroll up to the top of the page, and then click on the “Submit” button to update the Genome Browser display
Figure 7  Click on the “+” button underneath the “Minus” column label in the “Select subtracks by strand and stage” section (red arrow) to show the minus strand RAMPAGE data for all of the available developmental stages.

Examination of the RAMPAGE data for all of the available developmental stages shows a more complex picture of transcription initiation for the sev gene. Examine Figure 8 and your genome browser, and note that some developmental stages have a RAMPAGE signal at a different genomic position, or have an additional RAMPAGE signal besides the TSS annotated by FlyBase. For several developmental stages, there is no RAMPAGE signal.

Figure 8  RAMPAGE read density for different stages of development.
Q8. Do you think there is a TSS for each RAMPAGE signal (e.g., examine the RAMPAGE signal for 9-hour embryos)? Why or why not?

To compare the TSS identified by RAMPAGE with the mRNA expression levels:

- Under “RNA Seq Tracks”
  - Click on the blue “modENCODE RNA-Seq (Development) (R5)” link
    - Scroll down to the “Select subtracks by strand and stage” section
    - Click on the “-” button at the top left corner to unselect all the datasets
    - Click on the “+” button underneath the “Minus” column label
    - Scroll up to the top of the page, and then click on the “Submit” button to update the Genome Browser display

Q9. How many developmental stages show no RAMPAGE signal? How does the lack of RAMPAGE read density relate to the RNA expression levels of the sev gene during these developmental stages?

Q10. Based on the analysis of the CAGE and RAMPAGE data, do you think all of the developmental stages use the same TSS?

RNA POL II CHIP-SEQ

Chromatin immunoprecipitation followed by high throughput sequencing (ChIP-Seq) is a valuable method that can be used to identify DNA binding sites for proteins such as transcription factors. The first step of chromatin immunoprecipitation (ChIP) is to treat tissues or cells with formaldehyde to covalently crosslink DNA with DNA-binding proteins. The cross-linked DNA is then sheared so that the fragments are approximately 100–300 bp in size. Antibodies that recognize the protein of interest are added, and form a
precipitate that can be separated from the rest of the proteins and DNA. The crosslinked DNA is treated to remove the protein of interest, then sequenced (hence the name ChIP-Seq) and mapped to the genome assembly.

This technique can be used to identify modifications on histones as well, since the antibodies used for immunoprecipitation can recognize post-translational modifications including methylation, acetylation, and phosphorylation, to the histone proteins.

More recently, X-ChIP-Seq has been used to identify where RNA polymerase II (RNA Pol II) is enriched along the gene\(^2\) (see [https://www.illumina.com/science/sequencing-method-explorer/kits-and-arrays/x-chip.html](https://www.illumina.com/science/sequencing-method-explorer/kits-and-arrays/x-chip.html) for a figure that illustrates this method). DNA is crosslinked to histone proteins, then nuclease is added to cleave the linker DNA between nucleosomes. An antibody against one of the proteins in the RNA Pol II complex is used to immunoprecipitate DNA bound by the polymerase. The DNA is then sequenced and mapped to the genome, as with other types of ChIP-Seq.

The results show the genomic regions that are enriched in RNA Pol II, which usually corresponds to the approximate location of TSSs. The RNA Pol II X-ChIP-Seq data can also be used to identify genes whose expression is regulated by RNA Pol II pausing, where RNA Pol II is paused between 30–60nt downstream of the TSS after the initiation of transcription.

To examine the X-ChIP-Seq data for the sev gene more closely:

- Change the “enter position or search terms” field to “chrX:11,086,192-11,086,362” and then click “go”.
- Click on the “hide all” button to hide all the evidence tracks
- Under “Mapping and Sequencing Tracks”
  - Base Position: full
  - Click on the blue “Short Match” link
    - Verify the “TCAKTY” sequence is in the “Short (2-30 base) sequence” text box
    - Change the “Display mode” to “pack”
    - Click on the “Submit” button
- Under “Gene and Gene Prediction Tracks”
  - FlyBase Genes: pack
- Under “ChIP Seq Tracks”
  - RNA PolII X-ChIP-Seq: full
- Under “RNA Seq Tracks”
Click on the blue “Combined modENCODE RNA-Seq (Development) (R5)” link
- Change the “Display mode” field to “full”
- Scroll down to the “List subtracks” section
- Uncheck the box for the “modENCODE RNA-Seq (Plus)” track
- Scroll up to the top of the page, and then click on the “Submit” button to update the Genome Browser display

Figure 9 Enrichment of RNA Pol II near the 5’ end of the sev gene, and the combined RNA-Seq data on the minus strand.

Q11. According to the X-ChIP-Seq Log Likelihood Enrichment track (Figure 9), what are the coordinates of the region that is enriched in RNA Pol II?

INVESTIGATION 2: IDENTIFICATION OF TSS USING SEQUENCE CONSERVATION DATA

CONSERVATION

Comparison between closely related species of Drosophila can be used to identify conserved sequences near the TSS. In general, we expect the coding regions of a gene will be the most highly conserved (because it is under the strongest selective pressure), followed by the untranslated regions, and regulatory motifs (e.g., transcription factor
binding sites). In Investigation 2, you will learn how to configure the genome browser in order to view sequences from 26 species of Drosophila. The sequences have been "aligned"; that is, the sequences are shown in rows to make it easy to compare them.

Sophisticated computer software programs called LAST and ROAST are used to generate the multiple sequence alignments for a collection of species at different evolutionary distances from *D. melanogaster*. The genome assemblies for 26 Drosophila genomes are compared against each other using LAST. The program ROAST analyzes these pairwise alignments to construct a multiple sequence alignment, using *D. melanogaster* as the reference. You can read more about these programs by clicking on the blue “Drosophila Conservation (26 Species)” link under the “Comparative Genomics” section.

To examine the Conservation track for the *sev* gene more closely:

- Under “Expression and Regulation”
  - RNA Pol II X-ChIP-Seq: hide
- Under “RNA Seq Tracks”
  - Combined modENCODE RNA-Seq (Development) (R5): hide
- Under “Comparative Genomics”
  - Drosophila Conservation (26 Species): full

- Change the “enter position or search terms” field to “chrX:11,086,250-11,086,349” and then click “go”.

**Figure 10** (page 15) shows a multiple sequence alignment of the region near the TSS of the *sev* gene from 26 species. Note that the sequence near the TSS is highly conserved — the Initiator motif is conserved in *D. melanogaster* and 15 other *Drosophila* species.
Figure 10  Multiple sequence alignment of 26 Drosophila species with conservation scores from PhastCons and phyloP for the region surrounding the TSS of sev. (Inset) The Initiator motif in D. melanogaster is conserved in 15 other Drosophila species.

Q12. What is the sequence of the Initiator motif for the sev gene?

Q13. Based on the location of the conserved Initiator motif, what is the coordinate of the first nucleotide of exon 1?

REFERENCES
