Electrophoretic Mobility Shift Assays for RNA–Protein Complexes

Donald C. Rio

The electrophoretic mobility shift assay (EMSA), or gel mobility shift assay, is a popular and powerful technique for the detection of RNA–protein interactions. It relies on the fact that naked RNA has certain mobility on nondenaturing gels, but if the RNA is bound by protein, the mobility of the RNA is reduced. Therefore, the binding of protein results in a characteristic upward shift of the RNA on a gel, as monitored using radiolabeled RNA. For reasons that are not completely understood, most RNA–protein complexes—particularly those that result from high-affinity interactions—do not dissociate during the prolonged times required for electrophoretic separation. Because high-affinity interactions are more stable, it is often possible to identify specific interactions over a “background” of weak interactions. Accordingly, EMSAs can be performed using complex mixtures of proteins such as cell extracts. They can be used to investigate a wide range of RNA–protein interactions—from single protein-binding events to assembly of large complexes such as the spliceosome. EMSAs can also be useful for determining kinetic parameters (such as affinity constants) for RNA–protein interactions.

MATERIALS

It is essential that you consult the appropriate Material Safety Data Sheets and your institution’s Environmental Health and Safety Office for proper handling of equipment and hazardous material used in this protocol.

RECIPIES: Please see the end of this protocol for recipes indicated by <R>. Additional recipes can be found online at http://cshprotocols.cshlp.org/site/recipes.

Reagents

Acrylamide (40% stock)
Binding buffer for mobility shift assays <R>
Bisacrylamide (2% stock)
Bovine serum albumin (BSA)
Bromophenol blue
Glycerol
Heparin (50 mg/mL [w/v] in H2O) (Sigma-Aldrich H3393)
Loading buffer for mobility shift assays <R>
NaCl
Protein of interest (purified protein or cell extract)

See Preparation of Nuclear Extracts from HeLa Cells (Nilsen 2013).


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Cite this protocol as Cold Spring Harb Protoc; doi:10.1101/pdb.prot080721
RNA of interest (uniformly labeled and purified)


Splicing mix for mobility shift assays (4×) <R>
Total RNA or tRNA (10 mg/mL from yeast or Escherichia coli)
Tris (pH 7.6)
Tris/borate/EDTA (TBE) electrophoresis buffer (10×) <R>
Tris–glycine buffer (10×) <R>
Xylene cyanol

Equipment

Electrophoresis equipment

This should include a vertical gel apparatus; plates (17 × 14.7 × 0.15 cm), comb (with 0.75 cm-wide wells), and spacers (0.8 or 1.5 mm thick) for pouring gels; and a high-voltage power supply.

Ice

Incubator at appropriate temperature for binding (4°C–37°C)

Microcentrifuge tubes (1.5 mL)

Phosphorimager equipment or X-ray film and intensifying screen

Vacuum gel dryer

Whatman 3MM paper

METHOD

It is important to note that there are a large number of potential variations to gel-shift assays. Thus, the buffers, incubation times and temperatures, nonspecific competitors (see Discussion), and even gel conditions may vary depending on the complexes being studied. Below are example protocols for different types of RNA–protein interactions—from simple interactions (e.g., U1 snRNA and purified U1A protein) and small complexes (e.g., U1 snRNA and its associated proteins) to large multicomponent complexes (e.g., the spliceosome). These are readily adaptable to any situation.

Simple Interactions

1. Label the RNA of interest.

   • To label the RNA uniformly with [α-32P]NTPs, use in vitro transcription (see In Vitro Transcription of Labeled RNA: Synthesis, Capping, and Substitution [Nilsen and Rio 2012]).

   • For an oligoribonucleotide or a 100–200-nucleotide purified RNA, end-label using [32P]cytidine 3′,5′-bis(phosphate) or [γ-32P]ATP (see 3′-End Labeling of RNA with [5′-32P]Cytidine 3′,5′-Bis(Phosphate) and T4 RNA Ligase 1 [Nilsen 2014] and 5′-End Labeling of RNA with [γ-32P]ATP and T4 Polynucleotide Kinase [Rio 2014]).

2. Incubate the labeled RNA with purified protein under suitable binding conditions (e.g., binding buffer supplemented with 10 µg/mL BSA). For example, mix the following together on ice.

   Binding buffer (supplemented with 10 µg/mL BSA) 20 µL
   RNA (50,000 cpm) 1 µL
   Purified protein (serial dilutions from excess to less than equimolar) 4 µL

   Incubate the mixture for 30–60 min at room temperature, 30°C, or 37°C.
   
   It is often useful to include controls, such as inhibitors of the interaction and both specific and nonspecific RNA competitors.

3. After incubation, dilute the sample with 4 µL of loading buffer (without heparin) and load 10–20 µL of the mixture onto a native polyacrylamide gel. For a single protein (≏50,000 MW)
and a single RNA (~100 nucleotides long), start with a 0.8 mm-thick 6% acrylamide:bisacrylamide (30:1) gel made with Tris–glycine buffer or TBE. Prerun the gel for 30 min.

Try varying the pH of the gel and buffers. Vary the ratio of Tris to glycine until the optimal pH is found.
The percentage of gel and ratio of acrylamide:bisacrylamide may need to be varied to optimize separation of the bound and free RNA.
Do not overload the wells. For small wells (20-well comb), use a maximum volume of 5 µL/well.

4. Perform electrophoresis at 300 V for 3 h (or 13 V/cm), making sure that it does not heat up. If the interactions are suspected to be unstable, run the gel in a cold room.

5. After the gel is run, use a sheet of Whatman 3MM paper to pick up the gel. Dry the gel with vacuum at 80°C. Expose the gel to a phosphorimager or X-ray film.

Longer RNA Forming a Small Complex

1. Prepare a labeled RNA target by in vitro transcription (see In Vitro Transcription of Labeled RNA: Synthesis, Capping, and Substitution [Nilsen and Rio 2012]). Gel-purify the RNA to make sure that the substrate will be a single band on a gel.

2. Incubate the labeled RNA with purified protein under suitable binding conditions. For example, mix the following together on ice.

<table>
<thead>
<tr>
<th>Component</th>
<th>Volume</th>
</tr>
</thead>
<tbody>
<tr>
<td>Binding buffer</td>
<td>20 µL</td>
</tr>
<tr>
<td>RNA (50,000 cpm)</td>
<td>1 µL</td>
</tr>
<tr>
<td>Purified proteins or cell extract</td>
<td>1–4 µL</td>
</tr>
</tbody>
</table>

   Incubate for 30–60 min at the preferred temperature (4°C–37°C).

   It is often useful to include controls, such as inhibitors of the interaction and both specific and nonspecific RNA competitors.

3. Following incubation, dilute the reaction to 100 µL with buffer containing 500 mM NaCl, 10 mM Tris (pH 7.6), 5% glycerol, and 200 µg/mL heparin (in cases where cellular extract is used). Incubate for 10 min at room temperature.

   Heparin and high salt are included to resolve what could otherwise appear as a radioactive smear into discrete complexes on the gel. However, heparin also likely strips some proteins from the complexes. Total RNA or tRNA might also be added to improve resolution.

4. Remove 1 µL of the reaction and add it to 30 µL of loading buffer. Make sure that the glycerol concentration is at least 5% so that the sample does not float away.

5. Load this sample onto a 1.5 mm-thick, 20 cm-long 0.5 × TBE, 6% acrylamide:bisacrylamide (60:1) gel. Perform electrophoresis at 25 mA until the xylene cyanol is about two-thirds down the gel.

   The percentage of gel and ratio of acrylamide:bisacrylamide may need to be varied to optimize separation of the bound and free RNA.

   Do not overload the wells. For small wells (20-well comb), use a maximum volume of 5 µL/well.

6. Disassemble the gel, pick it up with Whatman 3MM paper, and dry it at 80°C. Expose the gel to a phosphorimager or X-ray film.

Large Multicomponent RNA–Protein Complexes

1. Prepare a native gel containing 3.75%–4.2% polyacrylamide and acrylamide:bisacrylamide (80:1).

2. Pour the solution into 17 × 14.7 × 0.15 cm plates, insert a comb with 0.75 cm-wide wells, and allow the gel to polymerize. (This usually takes at least 30 min.)

   The gel will be very jelly-like. Narrower wells, such as those in a 20-well comb, can also be used.

3. Mix the following components in a 1.5 mL microcentrifuge tube by pipetting up and down.
Nuclear extract (15–20 mg/mL) or extract dialysis buffer 5 µL
RNA (usually 25,000–50,000 cpm) 1 µL
Splicing mix (4×) 5 µL
H₂O 9 µL

Incubate for 30–60 min at 20°C–30°C.

It is often useful to include controls, such as inhibitors of the interaction and both specific and nonspecific RNA competitors.

4. Stop the reaction by placing the tubes on ice. Add heparin (titrate a range between 5 and 500 µg/mL) to reduce nonspecific binding.

5. Add 3 µL of 1% xylene cyanol and bromophenol blue to the reactions. Mix by pipetting up and down.

Alternatively, the dye can be added to a free lane on the gel to monitor electrophoresis and prevent possible interference with protein binding.

6. Prerun the gel for 20 min at no more than 6 W.

For large protein–RNA assemblies, such as pre-mRNA splicing complexes, it may be useful to try native agarose gels instead of polyacrylamide gels.

7. Load 5 µL of the binding reaction mixture from Step 5 onto the gel.

Do not overload the wells. For small wells (20-well comb), use a maximum volume of 5 µL/well.

8. Run the gel at no more than 6 W at 120–210 V maximum at room temperature until the dye is 1–2 cm from the bottom (for a 70-nucleotide RNA).

9. Transfer the gel onto Whatman 3MM filter paper and dry it for 30–60 min at 80°C on a vacuum gel dryer.

10. Expose the gel to X-ray film with an intensifying screen at −80°C.

DISCUSSION

EMSAs are quite versatile and can be used for a wide variety of purposes (see, e.g., Bowers et al. 2006; Das and Reed 1999; Konarska 1989). They can be valuable for determining if a protein of interest is present in a particular preparation of proteins (e.g., cytoplasmic or nuclear extract). In this case, a labeled synthetic RNA would be incubated with the protein preparation. Following nondenaturing gel electrophoresis, the labeled RNA is visualized by phosphorimaging or autoradiography, and the presence of shifted (or not) material is observed. If a gel shift is observed, it is then necessary to determine if the shift resulted from the binding of a protein that has a specific binding site in RNA.

To distinguish between specific and nonspecific binding, competition experiments are performed. If the shift is abolished in the presence of an excess of specific competitor (unlabeled RNA identical to the labeled RNA), but not by an excess of nonspecific competitor (unlabeled unrelated RNA or a mutant version of the labeled RNA), it can be concluded that the observed shift resulted from specific binding.

If specific binding is observed and one suspects that a specific protein is responsible and an antibody is available for that protein, it is possible to perform what is called supershift analysis. Here, the antibody is included in the initial binding incubation before electrophoresis. Binding of the antibody to a complex of interest with the RNA results in a further reduction in mobility of the complex (i.e., a supershift).

Other more complex questions can also be addressed. Mobility shift approaches have been extremely valuable in deciphering the ordered pathway of assembly of large (megadalton) complexes such as the spliceosome. In this case, assembly can be monitored in the presence or absence of an energy source (ATP) or a specific RNP (small nuclear ribonucleoprotein) or auxiliary factor(s). Such an approach is often the method of choice whenever an RNA–protein interaction (or more complex assembly) can be monitored in vitro.
Although conceptually straightforward, it is important to note that gel composition and percentage as well as electrophoresis conditions can have profound effects on the results obtained. Finally, it is worth noting that gel mobility shift assays can be used to quantitate various kinetic parameters of RNA–protein interactions (Setzer 1999; Goodrich and Kugel 2007), although, of course, pure proteins (or RNPs) are required for such applications.

**RECIPEs**

**Binding Buffer for Mobility Shift Assays**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Quantity (for 10 mL)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris (1 M; pH 8.0)</td>
<td>0.4 mL</td>
<td>40 mM</td>
</tr>
<tr>
<td>KCl (2 M)</td>
<td>0.15 mL</td>
<td>30 mM</td>
</tr>
<tr>
<td>MgCl$_2$ (1 M)</td>
<td>10 µL</td>
<td>1 mM</td>
</tr>
<tr>
<td>NP40 (10%, w/v)</td>
<td>10 µL</td>
<td>0.01% (w/v)</td>
</tr>
<tr>
<td>Dithiothreitol (DTT) (1 M)</td>
<td>10 µL</td>
<td>1 mM</td>
</tr>
<tr>
<td>H$_2$O or extract</td>
<td>9.42 mL</td>
<td></td>
</tr>
</tbody>
</table>

If the proteins or extract used for binding are not in glycerol, add 5% to the binding buffer. Store indefinitely at −20°C.

**Loading Buffer for Mobility Shift Assays**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Quantity (for 10 mL)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>KCl (2 M)</td>
<td>0.3 mL</td>
<td>60 mM</td>
</tr>
<tr>
<td>Tris (1 M; pH 7.6)</td>
<td>0.1 mL</td>
<td>10 mM</td>
</tr>
<tr>
<td>Glycerol (50%, w/v)</td>
<td>1 mL</td>
<td>5% (w/v)</td>
</tr>
<tr>
<td>Heparin (50 mg/mL)</td>
<td>40 µL</td>
<td>200 µg/mL</td>
</tr>
<tr>
<td>Xylene cyanol (0.1%, w/v)</td>
<td>1 mL</td>
<td>0.01% (w/v)</td>
</tr>
<tr>
<td>Bromophenol blue (0.1%, w/v)</td>
<td>1 mL</td>
<td>0.01% (w/v)</td>
</tr>
<tr>
<td>H$_2$O</td>
<td>6.56 mL</td>
<td></td>
</tr>
</tbody>
</table>

Store at 4°C. The amount of heparin may need to be adjusted or omitted.

**Splicing Mix for Mobility Shift Assays (4×)**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Quantity (for 100 µL)</th>
<th>Final concentration (4×)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Creatine phosphate (0.2 M)</td>
<td>10 µL</td>
<td>20 mM</td>
</tr>
<tr>
<td>HEPES/KOH (1 M; pH 7.6)</td>
<td>8 µL</td>
<td>80 mM</td>
</tr>
<tr>
<td>MgCl$_2$ (100 mM)</td>
<td>12 µL</td>
<td>12 mM</td>
</tr>
<tr>
<td>ATP (100 mM)</td>
<td>8 µL</td>
<td>8 mM</td>
</tr>
<tr>
<td>Dithiothreitol (100 mM)</td>
<td>40 µL</td>
<td>40 mM</td>
</tr>
<tr>
<td>H$_2$O</td>
<td>22 µL</td>
<td></td>
</tr>
</tbody>
</table>

Prepare fresh for each use. Keep on ice.

**TBE Electrophoresis Buffer (10×)**

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Quantity (for 1 L)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris base</td>
<td>121.1 g</td>
<td>1 M</td>
</tr>
<tr>
<td>Boric acid</td>
<td>61.8 g</td>
<td>1 M</td>
</tr>
<tr>
<td>EDTA (disodium salt)</td>
<td>7.4 g</td>
<td>0.02 M</td>
</tr>
</tbody>
</table>

Prepare with RNase-free H$_2$O. Dilute 100 mL to 1 L to make gel running buffer. Store for up to 6 mo at room temperature.
### Tris–Glycine Buffer (10×)

<table>
<thead>
<tr>
<th>Reagent</th>
<th>Quantity (for 1 L)</th>
<th>Final concentration</th>
</tr>
</thead>
<tbody>
<tr>
<td>Tris base (1 M)</td>
<td>500 µL</td>
<td>0.5 M</td>
</tr>
<tr>
<td>Glycine (1 M)</td>
<td>500 µL</td>
<td>0.5 M</td>
</tr>
</tbody>
</table>

Store at 4°C. The pH should be ~8.8.

### REFERENCES


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Cold Spring Harb Protoc; doi: 10.1101/pdb.prot080721

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