Accelerating eCLIP Studies: Importance of Antibody Validation and Pre-Validated Antibodies for RNA Binding Proteins

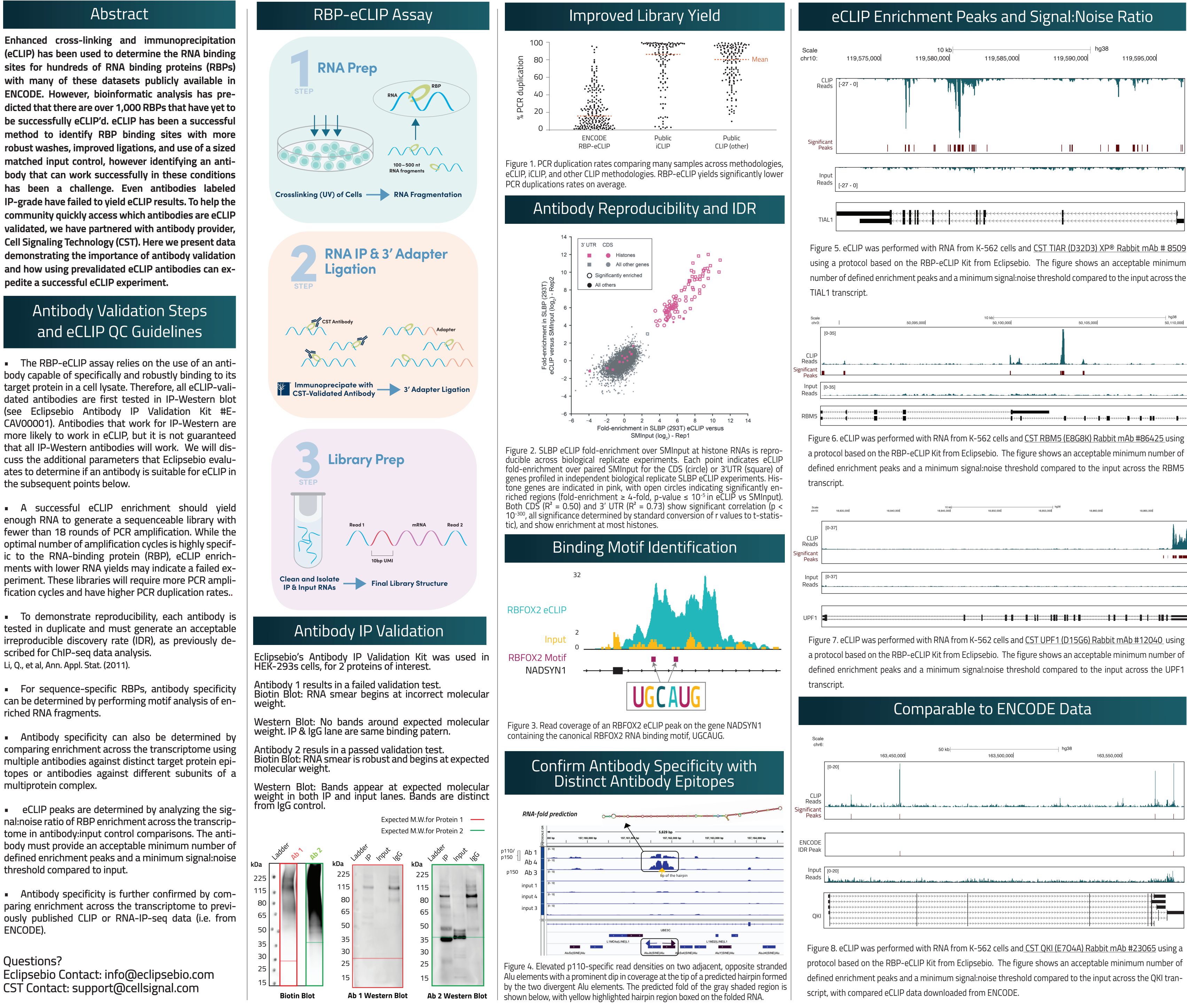
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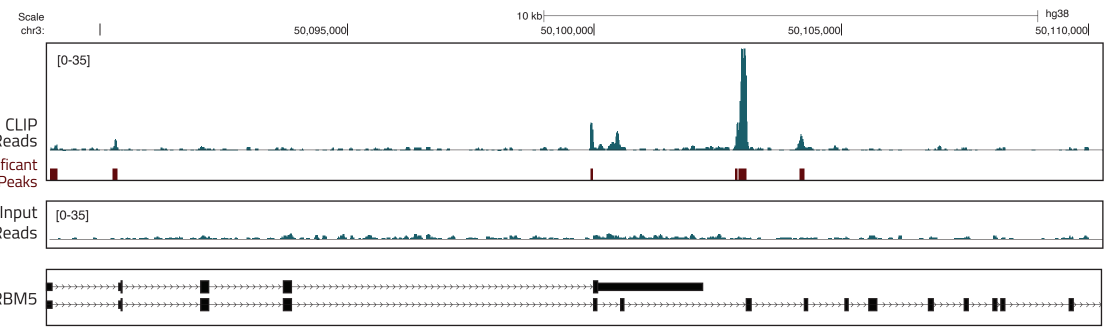
Enhanced cross-linking and immunoprecipitation (eCLIP) has been used to determine the RNA binding sites for hundreds of RNA binding proteins (RBPs) with many of these datasets publicly available in ENCODE. However, bioinformatic analysis has predicted that there are over 1,000 RBPs that have yet to be successfully eCLIP'd. eCLIP has been a successful method to identify RBP binding sites with more robust washes, improved ligations, and use of a sized matched input control, however identifying an antibody that can work successfully in these conditions has been a challenge. Even antibodies labeled IP-grade have failed to yield eCLIP results. To help the community quickly access which antibodies are eCLIP validated, we have partnered with antibody provider, Cell Signaling Technology (CST). Here we present data demonstrating the importance of antibody validation and how using prevalidated eCLIP antibodies can expedite a successful eCLIP experiment.



• The RBP-eCLIP assay relies on the use of an antibody capable of specifically and robustly binding to its target protein in a cell lysate. Therefore, all eCLIP-validated antibodies are first tested in IP-Western blot (see Eclipsebio Antibody IP Validation Kit #E-CAV00001). Antibodies that work for IP-Western are more likely to work in eCLIP, but it is not guaranteed that all IP-Western antibodies will work. We will discuss the additional parameters that Eclipsebio evaluates to determine if an antibody is suitable for eCLIP in the subsequent points below.

• A successful eCLIP enrichment should yield enough RNA to generate a sequenceable library with fewer than 18 rounds of PCR amplification. While the optimal number of amplification cycles is highly specific to the RNA-binding protein (RBP), eCLIP enrichments with lower RNA yields may indicate a failed experiment. These libraries will require more PCR amplification cycles and have higher PCR duplication rates.

• To demonstrate reproducibility, each antibody is tested in duplicate and must generate an acceptable irreproducible discovery rate (IDR), as previously de-



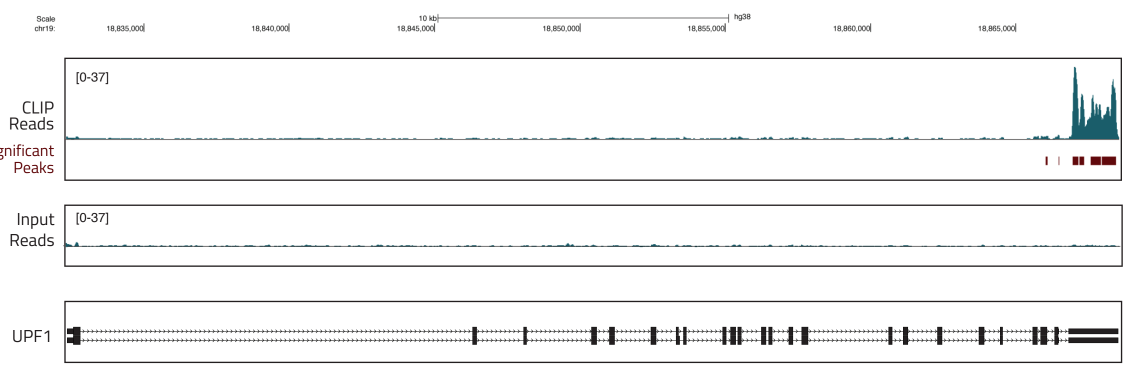


Figure 7. eCLIP was performed with RNA from K-562 cells and <u>CST UPF1 (D15G6) Rabbit mAb #12040</u> using