

# Robust transcriptome-wide identification of RNA binding protein targets

## Introduction

RNA binding proteins (RBPs) bind to RNAs through recognition of sequence and structural motifs to regulate RNA function in a cell-type, condition-specific, or temporal manner. Recent studies have estimated that there are over 1500 RBPs in the human genome and these RBPs play an integral role modulating RNA stability and function throughout the RNA life cycle. Mutations in RBPs have been linked to cancer, Amyotrophic Lateral Sclerosis (ALS), and numerous other diseases.

Enhanced crosslinking and immuno-precipitation followed by high-throughput sequencing (eCLIP) was developed to provide a robust and reproducible framework to map RBP binding sites on RNAs transcriptome-wide. Eclipsebio has optimized the eCLIP technology developed at UCSD and described in the 2016 Nature Methods paper to improve the efficiency of converting immunoprecipitated RNA into high-throughput sequencing libraries.

### Highlights

#### Highly efficient library prep

Increases experimental success rates & decreases wasted sequencing due to PCR duplication

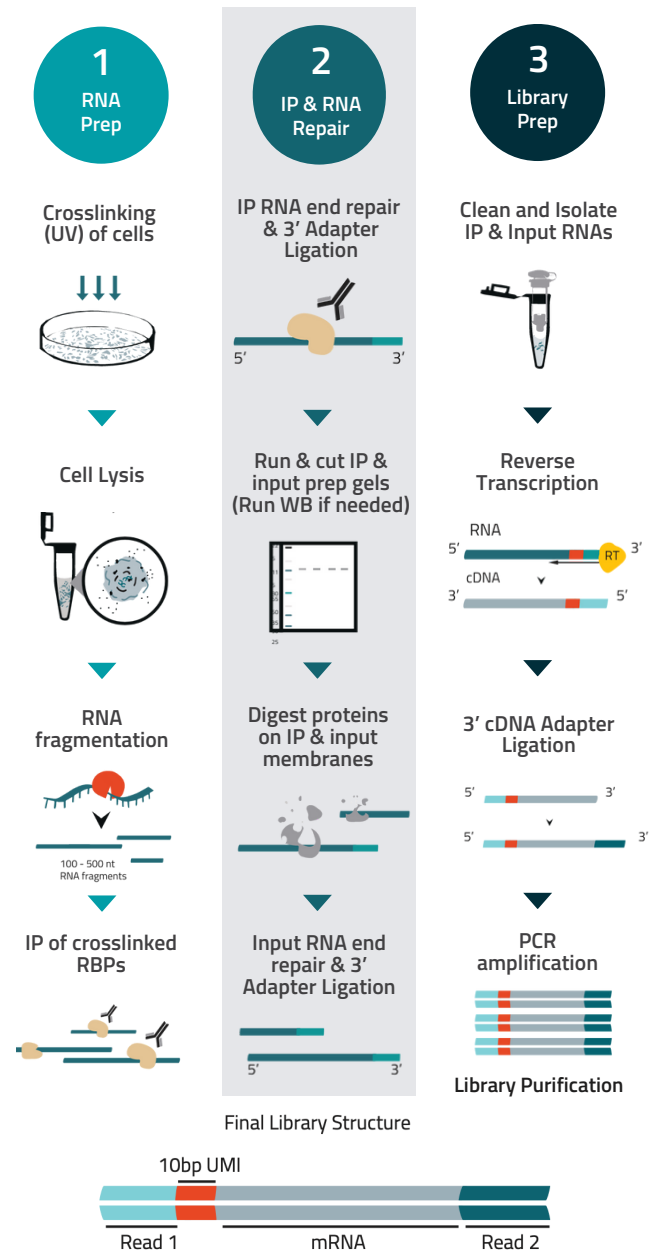
#### Transcriptome-wide targets

RBP binding sites discovered in all gene regions

#### RBP RNA binding motif identification

Validate known RNA binding motifs or discover novel RNA binding motifs

## RBP-eCLIP Workflow



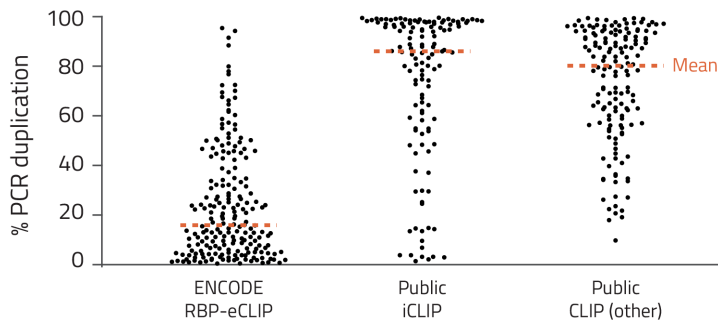
## Specifications

<b>Sample Input Range</b>	20M Cells or 80mgs Tissue
<b>Starting Material</b>	UV Crosslinked Cells
<b>Read Depth</b>	60M Reads/Sample
<b>Run Parameter</b>	SE100

**Figure 1.** RBP-RNA interactions are UV crosslinked. RNA is fragmented, and an RBP of interest is immunoprecipitated. After ligation of a 3' RNA adapter, IP material is run on denaturing protein gels and reverse transcribed to ssDNA (when a second adapter is ligated). PCR amplification is then used to obtain sufficient material for high-throughput sequencing.

## High efficiency library preparation with decreased PCR duplication

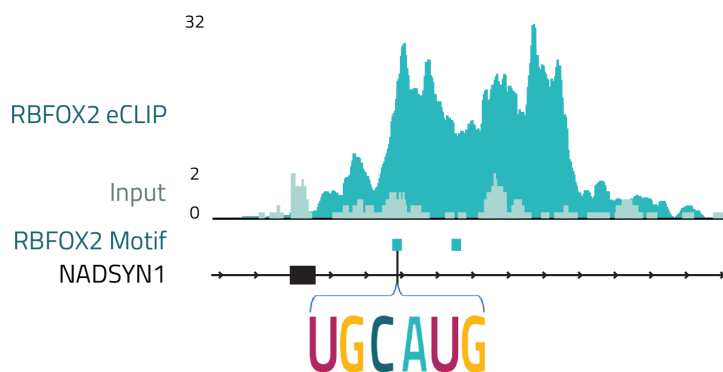
Optimization of the enzymatic steps of RBP-eCLIP improve library preparation by 1000-fold compared to other CLIP methods, thereby increasing experimental success rates and decreasing wasted sequencing due to PCR duplication.



**Figure 2.** PCR duplication rates comparing many samples across methodologies, eCLIP, iCLIP, and other CLIP methodologies. RBP-eCLIP yields significantly lower PCR duplications rates on average.

## RBP RNA binding motif identification

Validate known RNA binding motifs or discover novel RNA binding motifs



**Figure 3.** Read coverage of an RBFOX2 eCLIP peak on the gene NADSYN1 containing the canonical RBFOX2 RNA binding motif, UGCAUG

## Ordering Information

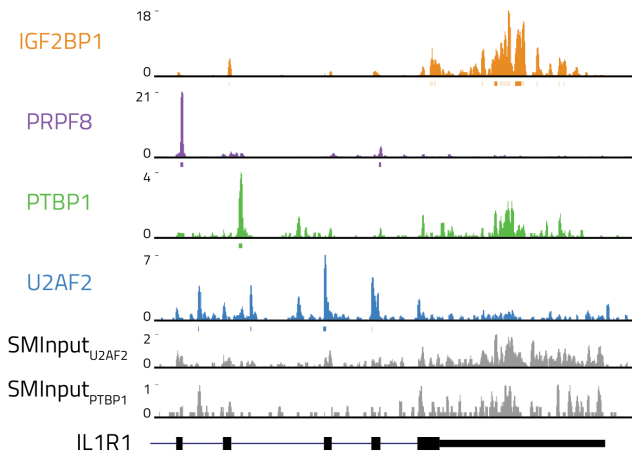
More information about RBP-eCLIP services online at [eclipsebio.com](http://eclipsebio.com) or contact us at [info@eclipsebio.com](mailto:info@eclipsebio.com).

## References

Van Nostrand EL, et. al. Robust transcriptome-wide discovery of RNA-binding protein binding sites with enhanced CLIP (eCLIP). *Nat Methods*. 2016 Jun;13(6):508-14

## Transcriptome-wide identification of RNA targets

RBP-eCLIP identifies RBP target binding sites across all genic regions: exons, introns and untranslated regions (UTRs), and in both coding and non-coding RNAs, including lincRNAs, microRNAs, and retrotransposons.



**Figure 4.** Read coverage showing peaks of reads representing binding sites for RBPs LIN28, IGF2BP1, PRPF8, PTBP1, and U2AF2 on the IL1R1 gene transcript.

## RBP-eCLIP Kit

An 8 sample kit for robust transcriptome-wide identification of your RNA-binding protein targets.



More information about RBP-eCLIP kit online at [eclipsebio.com](http://eclipsebio.com) or contact us at [info@eclipsebio.com](mailto:info@eclipsebio.com).