

Characterizing Novel Cyclophilin A-RNA

Interactions

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Abstract

RNA-binding proteins are a diverse and essential class of proteins with critical roles in key processes such as gene regulation, mRNA processing, and cell metabolism^{1,2}. While hundreds of RNA binding proteins (RBPs) have functions relating to RNA executed by well characterized RNA binding domains, a recent set of mRNA pulldown studies has uncovered a rich set of unexpected RBPs with no known binding domains and functions that do not obviously require RNA²⁻⁴. Among these putative RNA binders are proteins belonging to the cyclophilin family, proteins that isomerize the amino acid proline from the lower energy *trans* conformation to the higher energy *cis* conformation⁵.

Cyclophilin A (CypA), a highly conserved cyclophilin comprising a single isomerase domain⁶, was among the identified novel RBPs²⁻⁴. CypA is ubiquitous in the cell, as it abundant in the nucleus and cytoplasm as well as being secreted^{6,7}. It plays an essential role in protein folding and trafficking, cell signaling, and immune system response^{6,7} and has recently been shown to interact with key transcription factors to mediate their transcriptional activity^{8,9}. CypA is also biomedically relevant as the target of the immunosuppressive drug Cyclosporin A and is overexpressed in various cancers⁷. In addition to the mRNA studies, CypA and homologues have been implicated as RNA binders in several studies¹⁰⁻¹² though its isomerase activity does not seem to require RNA. It is thus an exciting model to study novel RNA-protein interactions. Here, I take a two-pronged approach to investigate CypA's RNA interactome: *in vitro* binding with previously implicated RNAs and *in vivo* RNA immunoprecipitation with subsequent bioinformatic analysis to identify relevant endogenous RNA classes that CypA may bind. The *in vitro* results suggest that CypA binds RNA with some specificity, as certain RNAs

interacted with CypA while others did not. The *in vivo* results differentiate a set of statistically significant enriched genes from a set of depleted genes, suggesting that CypA interacts sets of transcribed genes in the cell whose functional groups relate to known CypA activity.

Introduction and Background

RNA-binding proteins are essential in diverse cellular processes, ranging from gene expression and regulation to immune system response, governing such activities as mRNA splicing, translational regulation, and RNA stability, among others¹⁻⁴. Previous research has identified prevalent canonical RNA binding domains such as the OB fold¹³, RNA recognition motifs (RRMs)¹⁴, Zinc-finger domains¹⁵, and RGG and RG domains¹⁶. To understand the breadth of protein-RNA recognition, several unbiased studies have been done to characterize the mRNA-proteome of human cells to gain further insight into this important class of proteins²⁻⁴. In these studies, human cells were cross-linked with UV light to covalently link interacting molecules. All of the mRNA in the cells was then pulled down using a polyT tag that recognized the polyA tails of the mRNA and the attached proteins were identified using mass spectrometry²⁻⁴ (**Figure 1**). Interestingly, over 50% of the proteins identified using this unbiased strategy had no previously characterized RNA-binding domain²⁻⁴. Many of the identified proteins were metabolic enzymes whose functions do not apparently require RNA²⁻⁴. This led to exciting questions about RNA-protein interactions: how is RNA binding these unexpected proteins, what types of RNA molecules bind, and how does RNA interaction

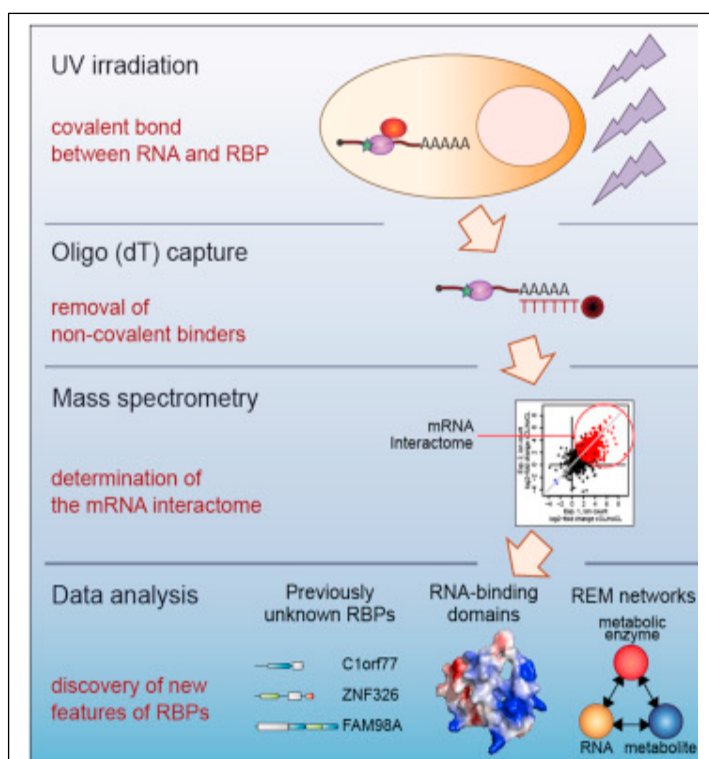
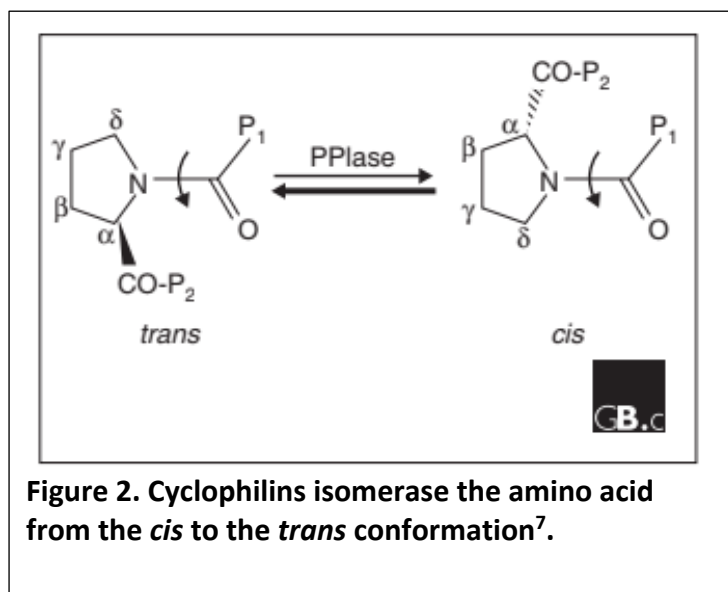


Figure 1. Identification of mRNA Binding Proteins. Study by Castello et al showing workflow. UV crosslinking, pulldown with polyA probe, identification of proteins via mass spectrometry,

contribute to, facilitate, or regulate these proteins' functions? Our lab chose the cyclophilin family of proteins to address these questions, as several were identified among the mRNA associated enzymes²⁻⁴.

The cyclophilin family is an essential group of proteins structurally conserved in diverse organisms across all domains of life^{5,6}. There are 17 human cyclophilins with various (some yet unknown) functions⁵, but all include a cyclophilin-like domain (CLD) of 109 amino acids that isomerizes the amino acid proline from the *trans* to the *cis* conformation^{5,6} (**Figure 2**⁷).

Due to its partial double-bond nature, the peptide bond of proline can exist in two conformations: the *trans* state, with amino acid side chains 180 degrees opposite, and the *cis* state, with adjacent side chains. Although both conformations exist in the cell, the *trans* conformation is slightly more energetically favorable. Cyclophilins stabilize the transition state for this isomerization and are thus critical in protein folding and assembly^{5,6}. Cyclophilins also play a role in cell signaling, such as CypA in the regulation of a kinase involved in T-cell activation⁷.



Some cyclophilins, such as CypE, are known to have roles in mRNA processing and splicing and contain canonical RRM (RNA recognition motifs) in addition to a CLD⁵, so it is not surprising that they were implicated in the transcriptome-bound proteome studies. However, other cyclophilins which contain only the CLD, specifically CypA and CypB, were also

implicated²⁻⁴. Experiments conducted previously in our lab show that inclusion of the CLD of CypE increases its RNA binding affinity compared to its RRM alone¹⁰. RNA was also shown to increase CypE and CypA's isomerase activity¹⁰. The cyclophilin-like domain, then, is an exciting model to explore as a non-canonical RNA binder.

Cyp A was chosen as a particularly interesting target as an essential, archetypal cyclophilin. It is a small (18kDa, 174 amino acids) protein comprising a single CLD.⁵⁻⁷ It is particularly ubiquitous— CypA is present in all mammalian tissues, found in the cytosol and nucleus, and is secreted⁷. In fact, it makes up about 0.1% of all cytosolic proteins⁶. Due to its high concentration in the cell, targeting it in *in vivo* studies is convenient: induced expression is not necessary— one can readily work with the endogenous quantities. CypA has been shown to mediate the activity of key transcription factors, including YY1⁸, NF-κB⁹, and the transcriptional circadian rhythm protein BMAL1.¹⁸ Isomerization by CypA and other proline isomerases of a key proline bond in BMAL1 causes a conformation change that leads to BMAL1 interaction with different activators and repressors.¹⁸ CypA is also involved in cell growth and is overexpressed in lung, breast, and liver cancer, among others⁷. Besides indicative of cancer, CypA is biomedically relevant as the target for the immuno-suppressive drug cyclosporin A.⁵⁻⁷ Cyclosporin A binds to CypA to form a ternary complex with calcineurin that blocks calcineurin's immune system activation⁵⁻⁷.

Previous RNA-CypA Interactions

As mentioned above, CypA's rate of isomerization has been shown to increase with the addition of a random RNA oligos in experimental assays¹⁰, thus implying functional significance

of the CypA-RNA interaction. CypA and its yeast homologue Cpr1, which shares 65% sequence identity with CypA, have also been shown to interact with the genomic RNA of the tomato bushy stunt virus (TBSV) to inhibit viral replication¹¹, and fungal *PiCypA* has also been shown to directly bind RNA¹². CypA is thus an evidentially implicated, medically relevant, and cellularly essential protein whose RNA binding activities may provide an unappreciated level of cellular regulation.

Here, I describe my work characterizing CypA-RNA interactions using biochemical and bioinformatic approaches to identify transcribed genes and RNA sequences CypA binds. I took a two-pronged approach: cross-linking and immunoprecipitation *in vivo*, followed by RNA sequencing and bioinformatic analysis, and Fluorescent Anisotropy (FA) binding assays with previously implicated RNA *in vitro*. These two complementary investigations provide good evidence that CypA binds RNA both *in vivo* and *in vitro*, opening exciting new avenues for future research.

Materials and Methods

In vitro

Protein Expression: Nickolaus Lammer, a graduate student in the Wuttke lab, expressed protein in transformed *Escherichia coli* (*E. coli*) cells prior to my joining the lab. CypA was expressed with a His-SUMO tag at its N-terminus in a pET28b plasmid. The plasmid was transformed into *E. Coli* cells, which grew at 37°C (with 50 g/ml kanamycin) in 2-X-Y-T media, then shocked on ice for 40 minutes. About 1 mL of 1 M of Isopropyl -D-thiogalactopyranoside was added per liter of culture to induce protein expression. Cultures were grown overnight at 18 °C. Cells were centrifuged at 5000 RCF to harvest and pellets stored at –20 °C.

Protein Purification: For the purification, cell pellets were resuspended in lysis buffer (50 mM Tris pH 8.5, 1000 mM NaCl, 10 mM imidazole pH 8.3, 10% glycerol, 0.1% Triton 100X) with an **EDTA**-free protease inhibitor. Lysis of cells was performed with a Misonix Sonicator 3000 with a half inch tip (15 seconds on, 15 seconds off, 10 to 12 times at 110 W) and then centrifuged for 30 minutes at 15,000 RCF. Lysate was poured over nickel-NTA beads and rocked at 4°C for 1 hour, then loaded onto a flow column. The column was washed three times with 15 mL of lysis buffer, then the protein was eluted twice with 15 mL of lysis buffer with 350 mM imidazole added (flow through collected each time). The protease Ulp1 was added to collected elution to cleave the His-tag^E. This solution was dialyzed overnight with SpectrumSpectra/Por Dialysis tubing with a MW cutoff of 6-8 kDa (Thermofisher) in a 4 L container of dialysis buffer (50 mM Tris pH 8.3, 135 mM KCl, 15 mM NaCl, 10 % glycerol). Cleaved, dialyzed solution was poured over the nickel column to remove uncleaved His-tagged proteins and the flow through containing CypA collected. The flow through was concentrated at

4 °C on a 5K MWCO concentrator to a volume of about 2 mL. It was then run through the G75 gel filtration column and fractions containing CypA were collected and pooled. Pooled fractions were again concentrated to about ~800 uM, aliquoted into 15 20 uL volumes, flash frozen in liquid nitrogen and stored at -70 °C. Protein yield was ~4.32 mg/L

RNA transcription: A double stranded 633 nt DNA gBlock was ordered from IDT (Integrated DNA Technologies, **Figure 3**) containing the sequence from the genome of the Tomato Bushy Stunt Virus. In addition, primers designed to segment the DNA into 5 sections with added T7 transcriptase recruiting sequences were ordered.

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GGAAATTCTCCAGGATTTCTCGACCTAGTTCGTTTATCTGGTGACTTGCGCTACCGTTG
CTTTGCGTAGAGAATTTCTCTCCATAATTATTATCTTTAGTTGTGGGGTTTGAAGGTTG
GGTCTACCTTTCGGGGGGGATAAATTGTAACCTCCAACAACAAGCGACATGTCTAGAA
GAAACGGGAAGCTCGCTCGCACTCCACAACCTACCAAAGGAGCCTTTGGACGTCTTC
CCCGTTCAGGAAAGCGGTTTGTGAGAAGTTGGGGTAGCCCACCGACTTGGGTATGA
TGGGTTTCTGTCATACTACAGCGGTGCGAAACTCCGTACTTACACACGAGCCGTGGAG
AGTCTGCATATCACACCTGTCTCCGAGAGGGATAGTCACTTGACTACCTTCGTA AAAAG
CAGAGA ACTGCAGAGCGAGTAAGACAGACTCTTCAGTCTGAGTTTGTGGAGATGAGT
GTAAATCTGGCATAGCATA CAGGTTACTCTTGTTGGGTATTCTGTTTACGAAAGTTA
GGTGTCACTTGTGGAAGCGGACCCAGACACGGTTGATCTCACCTTCGGGGGGGGCTA
TAGAGATCGCTGGAAGCACTACCGGACAACCGGAACATTGCAGAAATGCAGCCC
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Figure 3. Full Length DI-72 DNA gblock. DNA sequence corresponding to the RNA sequence of the viral DI-72 RNA

First, a PCR (polymerase chain reaction) was run to amplify DNA segments before transcription. The reactions were done in volumes of 50 uL (1x concentrated Taq buffer from 10x concentrated stock (15 mM MgCl₂, 500 mM KCl, 100 mM Tris-HCl pH 8.3), 200 uM dNTPs, 0.5 uM forward primer, 0.5 uM reverse primer, ~250 ng template DNA, 1 unit/50 uL Phusion DNA Polymerase, nuclease free water up to 50 uL). DNA was melted at 98 °C for 15 seconds, annealed at 52 °C

for 30 seconds (primers annealed to template), and elongated at 72 °C for 30 seconds. This sequence was repeated 25 times.

After amplification, segmented DNA was transcribed in 200 uL volumes (1X Transcription Buffer as used by Milligan et al.^F, 24 mM MgCl₂, 4 mM dATP, dGTP, dUTP, dCTP, 40 uL template from PCR, 97.6 uL nuclease free water, 10 mM DTT, 2 uL PPIase, 4 uL T7 polymerase). Reactions were left in the water bath for 2 hours. RNA purification was done by running a urea gel (7 M Urea, 8% Polyacrylamide) and cutting out RNA bands after visualization via UV shadowing.

Primer	Melting Temperature	Sequences
FWD1	57 °C	aaattctccaggatttctcgacc
REV1-T7	57-58 °C	TAATACGACTCACTATAGGcatgtcgcttgtttgttgaag
FWD2	59 °C	tagaagaaacgggaagctcgc
REV2-T7	59 °C	TAATACGACTCACTATAGGatcataccaagtctgggtggg
FWD3	58 °C	gggtttctgtcactactacagcg
REV3-T7	59 °C	TAATACGACTCACTATAGGttctctgcttttacgaaggtagtcaag
FWD4	58 °C	agcgagtaagacagactcttcag
REV4-T7	57 °C	TAATACGACTCACTATAGGaccaacaagagtaacctgtatg
FWD5	57 °C	attcctgtttacgaaagtttaggtgtc
REV5-T7	58 °C	TAATACGACTCACTATAGGctgcatttctgcaatgttccg

Figure 4. Sequences of DNA primers used to transcribe Minus Strand RNA sequences from DI-72 DNA gBlock. The lowercase letters are sections of the larger DI-72 sequence and the capital letters are transcription enzyme (T7) recruitment sequences. Shown are primer names, melting temperature, and sequence.

RNA 3'-end labeling with fluorescein-5-thiosemicarbazide (FTSC): To prepare transcribed RNA for fluorescence anisotropy binding assays, they were labelled at the 3' end with the fluorescent probe FTSC. For the oxidation reaction, 350 pmol RNA were added to 20 mM Sodium Periodate (NaIO_4) in a volume of 50 μL (nuclease free water up to 50 μL) and incubated in the dark at room temperature for 20 minutes. 7.5 μL of 2 M KCl (0.25 M) were added and samples rested on ice for 10 minutes. Reactions were spun for 10 minutes at 16.1k RCF and supernatant transferred to a new tube. 1 μL glycogen and 140 μL of 100% ethanol were added and samples frozen overnight. Samples were then pelleted, washed with 70% ethanol, and resuspended in 50 μL labeling solution (100 mM Sodium acetate pH 5.2, 1.5 mM FTSC). They were thoroughly mixed and incubated for 1 hour at 37 degrees Celsius. Samples were spun down, supernatant transferred to a new tube, 125 μL of 100% ethanol added, and samples then frozen overnight. Next, they were washed 4 times in 70% ethanol (pelleted via centrifugation, resuspended in 70% ethanol) and resuspended in 30 μL of RNA folding buffer. They were then passed through a G-25 spin column and the flow through was collected in an opaque tube. Verification of proper length and labelling effectiveness was done by running a denaturing PAGE gel and a dilution experiment in which intensity of diluted, labelled RNA was measured by a fluorimeter.

Fluorescence Anisotropy: A Fluorescence Anisotropy (FA) binding experiment was conducted with previously expressed and purified CypA and labelled RNA. Experiments were conducted with a constant concentration of RNA (3-6 nM) and 20 titrations of CypA (starting concentration of 66.5 nM, halved 20 times) in 20 μL binding buffer (50mM NaCl, 20 mM Tris, 10 % glycerol).

CypA and RNA incubated in the 384-well plate for 1 hour at room temperature before total anisotropy was measured. Anisotropy curves were fit to a single-site binding equation, $A = B + \frac{[RNA.CypA]}{[CypA] + K_d}$, where A is anisotropy, B is a baseline background, [CypA] is the concentration of CypA, [RNA.CypA] is the concentration of RNA bound to CypA. This simplified equation assumes that the ligand concentration is in sufficiently in excess (in this case, this will hold if the K_d for the CypA is sufficiently greater than RNA concentration). This assumption is valid given our conditions: reported K_d values were about 100-fold higher than RNA concentrations.

In vivo

Crosslinking: Due to COVID training restrictions that precluded me from access to the Biochemistry cell culture facility, a graduate student in the Wuttke lab (Nickolaus Lammer) facilitated the first step of this project by growing and cross-linking human HeLa cells using UV light or formaldehyde. The HeLa cells were grown in DMEM (Dulbecco's Modified Eagle Medium) containing 10% FBS and 1% penicillin-streptomycin solution. Crosslinking was done at 10-12 million cells. To induce crosslinking with UV light, 400 mJ/cm² of 254 nm UV light was applied to plated HeLa cells. For formaldehyde samples, cells incubated with 0.1% formaldehyde in 1 X PBS for 10 minutes. Crosslinking was quenched by adding 795 uL of 2.5 M glycine allowing plates to sit for 5 minutes. Both UV and formaldehyde crosslinked cells were then scraped and centrifuged (5 minutes at 1000 RCF), resuspended in 1 mL of PBS, centrifuged again at 4000 RCF for 75 seconds, and flash frozen (after supernatant removal) in liquid nitrogen. Frozen pellets were stored at -70 °C.

CypA-RNA Immunoprecipitation: Cross-linked cells were lysed with a biorupter in ~500uL lysis buffer (50 mM tris, pH 8, 150 mM NaCl, 0.1% SDS, 1% triton X-100, 5 mM **EDTA**, 0.5% sodium deoxycholate, 0.5 mM DTT, 1x Halt protease inhibitor cocktail (PIC) from Thermofisher, 100 units/mL of RNaseout). Cell lysate was collected after centrifugation and diluted with an equal volume of binding/washing buffer (25 mM Tris, pH 7.5, 150 mM KCl, 0.5% NP-40, 5 mM **EDTA**, 0.5 mM DTT, 1x PIC, and 100 U/mL RNaseOUT). This mixture incubated with 33 uL of protein G Dynabeads (Invitrogen) equilibrated with washing buffer (without PIC and RNase out) for 45 minutes to remove nonspecific bead binders. After magnetically removing beads, a 50uL aliquot was taken as an input and frozen at -20 °C. Then, the supernatant was incubated with 4.4 ug of mouse derived IgG anti-CypA antibody for 2 hours at 4°C. To pull down the CypA antibody, 50 uL of Dynabeads (Invitrogen) equilibrated in binding/washing buffer was added and incubated for 1 hour at 4 °C. The beads were pulled to the side and supernatant removed. Beads were washed twice with binding/washing buffer and frozen at -20 °C.

RNA Isolation: First, 45 uL of RNase or nuclease free H₂O and 33 uL of 3x reverse-crosslinking buffer (3x PBS (phosphate buffered saline), 6% N-lauroyl sarcosine, 30 mM **EDTA**, 15 mM DTT) and 2 uL RNaseOUT was added to the beads. 36 uL of reverse-crosslinking buffer and 2 uL of RNaseOUT was added to the input samples. Next, 20 uL of proteinase K was added to each input and bead samples and heated at 42 °C for 1 hour and for another hour at 55 °C. To separate RNA from other soluble molecules, 1 mL of trizol and 1.080 mL trizol were added to the beads and inputs, respectively, and vortexed. 200 uL of chloroform were added to beads and 216 uL chloroform to input followed by about 20 seconds of vortex. After centrifuging for

15 minutes at 16k RCF (relative centrifugal force), the aqueous layer was removed and 500 μ L of isopropanol (to precipitate RNA) was added per 1 mL of trizol used. After addition of 1 μ L of glycoblue and placing samples on ice for 10 minutes, they were centrifuged at 16 RCF for 15 minutes and the supernatant was removed. Samples air dried for about 5 minutes, then 85 μ L of nuclease free H₂O was added to each sample. DNA was then digested by adding 10 μ L of 10x DNase 1 reaction buffer and 1 μ L of DNase 1 to each sample, and they were allowed to sit at room temp for 15 minutes. RNA was then purified using Qiagen RNeasy MinElute Cleanup Kit (74204) quick start protocol. RNA samples were frozen at -20°C.

Library Preparation: Libraries were prepared for sequencing with KAPA RNA HyperPrep Kit with RiboErase (HMR) for Illumina Platforms. This protocol consisted of removing rRNA via hybridization to complementary DNA oligonucleotides followed by addition of RNase H to remove rRNA-DNA duplexes and DNase to remove excess DNA oligos. RNA was then fragmented to desired library size (~300 nt) using heat and magnesium. Due to its catalytic activities, RNA is able capable of self-cleavage, which is facilitated by Mg²⁺ ions stabilizing RNA structures conducive to cleavage and accelerated by heat^C. After RNA fragmentation for 6 min at 85 °C, the first strand of cDNA was synthesized with random priming, followed by synthesis of the second strand, which incorporated dUTPs to mark the strand and a dAMP to the 3' end. UMI (unique molecular identifiers) and Illumina adapters were then ligated via 3' dTMPs. Adapter sequences record condition (Formaldehyde or UV input or pulldown) and allow ligation to the flow cell, while UMIs allow accurate counts of each unique RNA molecule. Libraries were then amplified (only the strands without dUTP was amplified so that the original RNA molecules

would be sequenced). Adapter-ligated libraries were sequenced with Illumina NovaSeq Paired End (150 cycles) sequencing. Six samples in total were sequenced (an input and pulldown for two formaldehyde conditions and an input and pulldown for one UV condition). For the first formaldehyde experiment, 22,917,431 reads were obtained for the input and 28,187,490 for the pulldown. For the second, 30,177,674 were obtained for the input and 76,393,313 for the pulldown. For the UV experiment, 31,017,183 reads were obtained for the input and 31,339,559 for the pulldown.

Data Analysis Pipeline: The data pipeline for sequencing data consisted of trimming adapters and low-quality reads with Trim Galore (Barbara Bioinformatics), removing duplicate reads using UMIs with a python script written by Taeyoung Hwang, a postdoc in the Rinn lab, aligning reads to the human genome (GRCh38) using the STAR aligner, and counting the number of reads (RNA molecules) per gene or genomic feature with the Rsubread function featureCounts. The pipeline was primarily developed by a Taeyoung Hwang, a postdoc in the Rinn Lab. Further analysis (calculation of fold change IP/input and input) was done through a combination of bioinformatic software (DESeq2²¹) and python scripts I wrote. Clustering of enriched gene data was accomplished using DAVID²² tools. Code is available on request.

Results and Discussion

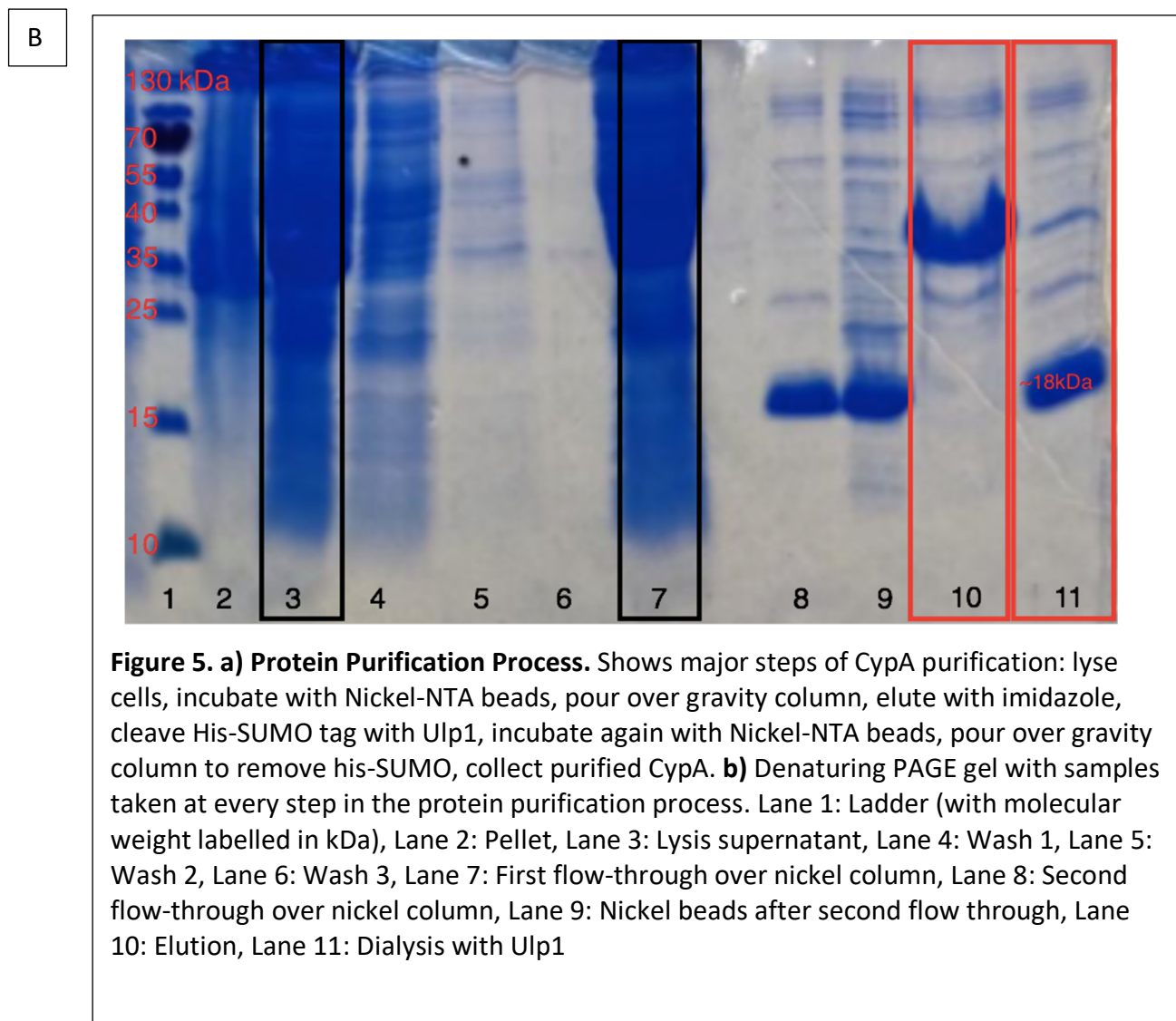
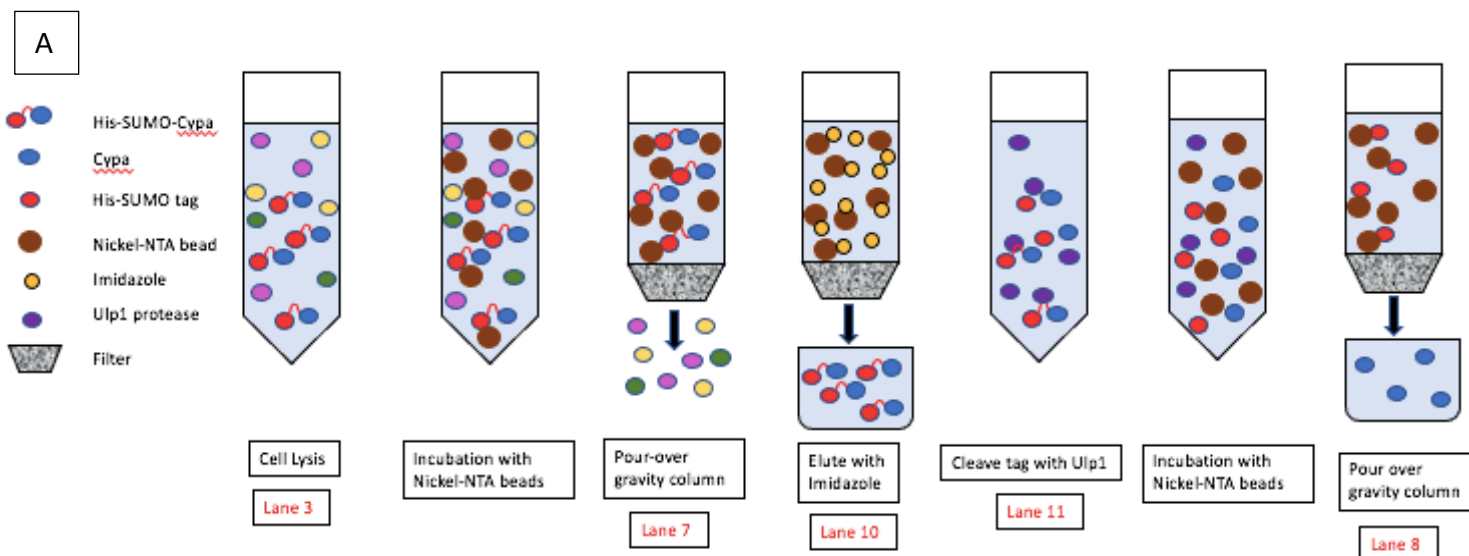
Fluorescence Anisotropy to Assess CypA DI-72 RNA Binding Affinity

CypA Expression and Purification

Before determining what RNAs CypA interacts with *in vivo*, I sought to ascertain if it binds RNA *in vitro* by expanding on previous studies that suggested RNA-CypA binding^{11,12}. Prior to conducting these *in vitro* binding assays, it was necessary to express and purify human CypA. CypA was expressed in *E. coli* cells with a His-SUMO tag at its N-terminus, which allowed for capture of the protein. The His-SUMO tag contained six histidine amino acids and a SUMO protein sequence. The histidines have a high affinity for binding metal ions²³, enabling the use of a commercially available nickel column to separate the tagged protein from other cellular components. The SUMO portion of the tag allows for removal of the tag from CypA after separation as it is recognized and cleaved by the protease Ulp1^B.

CypA purification consisted of lysing cells, pouring the supernatant (soluble cell parts) over a nickel column to which the histidine tag binds, elution of the protein and tag by washing with a high imidazole buffer (imidazole pushed the His-tag from the nickel column by competing for binding to charged metal beads), and cleavage of the tag from CypA with Ulp1 (**Figure 5A**). Clean-up of the tag was accomplished by another flow over the nickel column, and the resulting flow-through concentrated and further purified by gel filtration.

Figure 5B shows a 15% PAGE gel with samples taken at every step in the protein purification process. PAGE gels denature proteins and are thus used to separate them by size (larger proteins migrate more slowly and appear higher on the gel, while smaller proteins migrate more quickly and appear lower on the gel). Lanes 3 and 7 show the supernatant and



lanes contain all soluble cell proteins. While lane 7 should not include as high a concentration of His-SUMO-CypA (which is expected to stick to the column), the effect of this one protein construct's removal is indistinguishable from the signal that results from running all soluble cell components. Important to note are lanes 10 and 11. Lane 10 shows the elution (after the His-CypA has been competed off by the nickel column by imidazole). A thick band in the upper third of the lane can be seen. This is His-SUMO-CypA and indicates efficient protein expression and elution. Lane 11 shows a sample taken after dialysis, in which the His-SUMO tag was cleaved from CypA. In this lane, there is thick band that ran further than the tagged CypA, indicating successful cleavage from 32 kDa to 18 kDa. This band is slightly higher than the 15 kDa ladder marker, which is further reassuring as CypA's molecular weight is 18 kDa.

A second flow over the nickel column removed cleaved His-tags but did not purify the sample of unwanted higher molecular weight species (see Lane 8, Figure 3). These were removed by 120 mL size exclusion column, which separated the solution into 2 mL fractions of different molecular weight. **Figure 6** shows fractions collected at and surrounding CypA's molecular weight (18kDa). The band for CypA are clean and, again, run to slightly above the 15 kDa ladder mark, indicating a successful purification process.

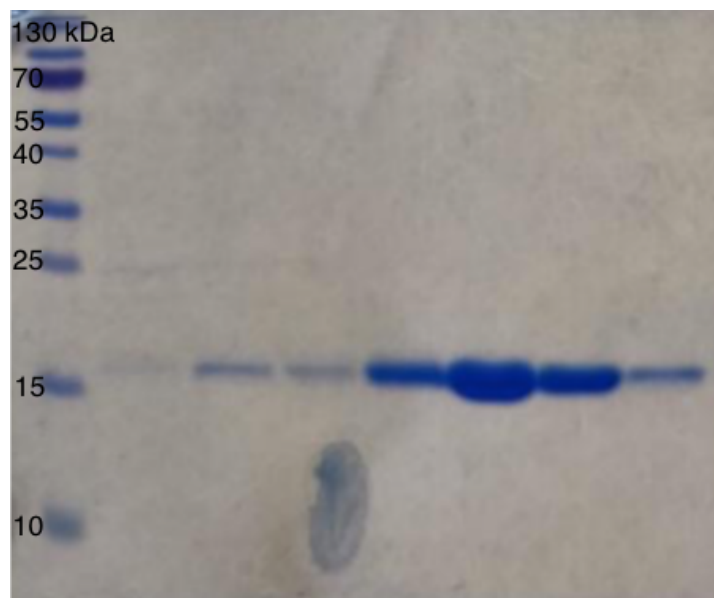


Figure 6. Purified CypA. Denaturing PAGE gel showing (left) ladder with labelled molecular weights and purified CypA fractions.

Selection and Transcription of RNA

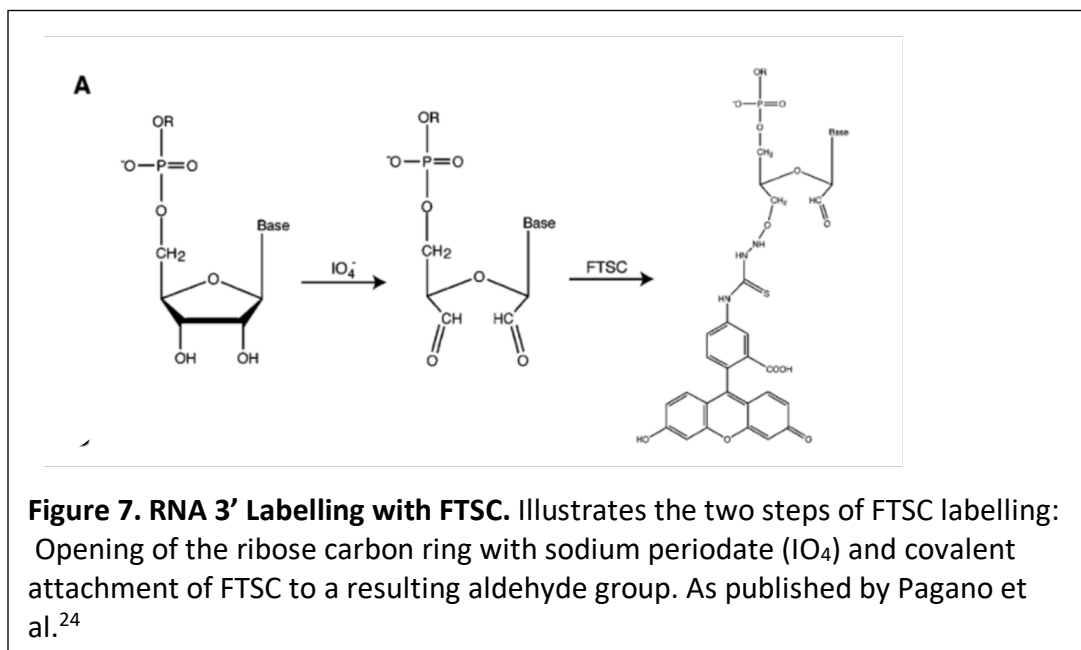
With a highly purified supply of CypA, I next obtained a suitable RNA molecule for *in vitro* investigation. Several previous publications provided RNA sequences that bound CypA or CypA homologues in plants and yeast^{11,12}. A particular relevant study in 2013 by Kovalev et al. showed that CypA inhibits the replication of the Tomato Bushy Stunt Virus capsid *in vitro*, plausibly by binding to the virus' genomic RNA¹¹.

Kovalev et al. tested CypA binding with a 633 nucleotide RNA sequence (referred to as DI-72) comprising several non-contiguous segments of the viral genome: the 5' and 3' ends, a template recognition and recruitment element from one of the protein coding sequences, and a replication enhancer¹¹. Both the plus strand (coding) and the minus strand (non-coding) were shown to bind CypA in a well-shift EMSA (Electromobility Shift Assay)¹¹. The minus strand bound more tightly than the plus strand, with an estimated dissociation constant (K_d) in the micromolar range¹¹. K_d measures the thermodynamic tendency of interacting molecules to fall apart and is calculated by dividing free concentrations of molecules by bound concentration at equilibrium: $\frac{[A][B]}{[AB]}$ where [A] and [B] are free concentration of molecules and [AB] is bound concentration. A lower K_d thus corresponds to tighter binding and higher affinity between two interactors. A well-shift experiment shows unbound RNA that runs into the gel lane and bound RNA that does not migrate from the well, thereby making it difficult to estimate an accurate K_d (intermediate concentrations between bound and unbound states are necessary). This estimated CypA-DI-72 K_d , then, was an exciting lead but warranted further investigation.

I thus obtained from the authors the full-length DI-72 sequence and ordered it as double-stranded DNA from which I could transcribe desired RNA segments. As mentioned

previously, the full-length molecule is 633 nt and I sought to hone-in on a smaller RNA sequence that CypA might bind. By designing specific DNA primers, I transcribed from the longer sequence five sections corresponding to the non-contiguous segments out of which they assembled the DI-72 molecule. I amplified the DNA, then transcribed five different sections (minus strand) with the corresponding primers. The five resulting oligos (referred to as Minus 1-5) were between 82 and 167 nucleotides long.

To prepare for a Fluorescence Anisotropy assay, the five RNA strands were labelled at their 3' end with FTSC (fluorescein-5-thiosemicarbazide), a fluorescent probe that absorbs light at 495 nm. The ligation of the fluorescent probe FTSC is a two-step process: first, the introduction of sodium periodate, which opens the carbon ring of ribose at the 3' end of the RNA molecule to form two reactive aldehyde groups and second, incubation with FTSC, which covalently attaches to the opened ring. **Figure 7**²⁴ illustrates the FTSC labelling process.



To check labelling efficiency, I ran the five RNA molecules on a Urea gel (denatures RNA) at the same concentration and excited the fluorophores with a gel imager using a fluorescent channel. **Figure 8** shows that the RNA molecules were effectively labelled as dark bands appear at the expected relative running distances. The Minus 1 strand was the longest of the five at 167 nucleotides, while Minus 4 strand was the shortest at 82 nucleotides. This was accurately reflected in the running distances. Strands 4 and 5 were not labelled with the same efficiency as the others (weak bands midway through the lane), making it necessary to use higher concentrations of these two strands in FA trials.

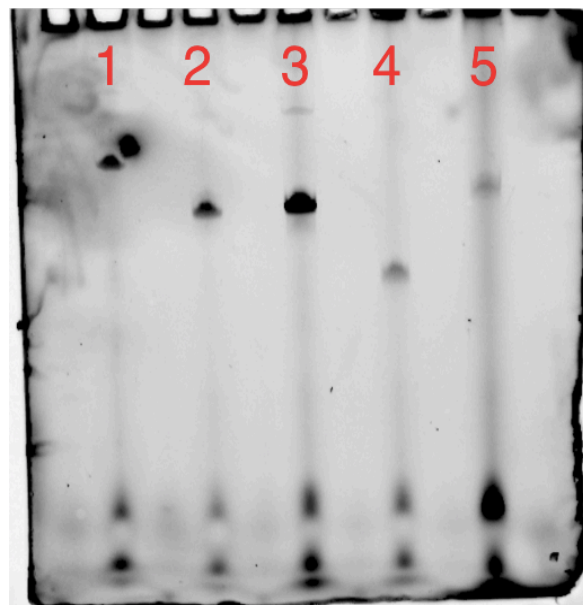
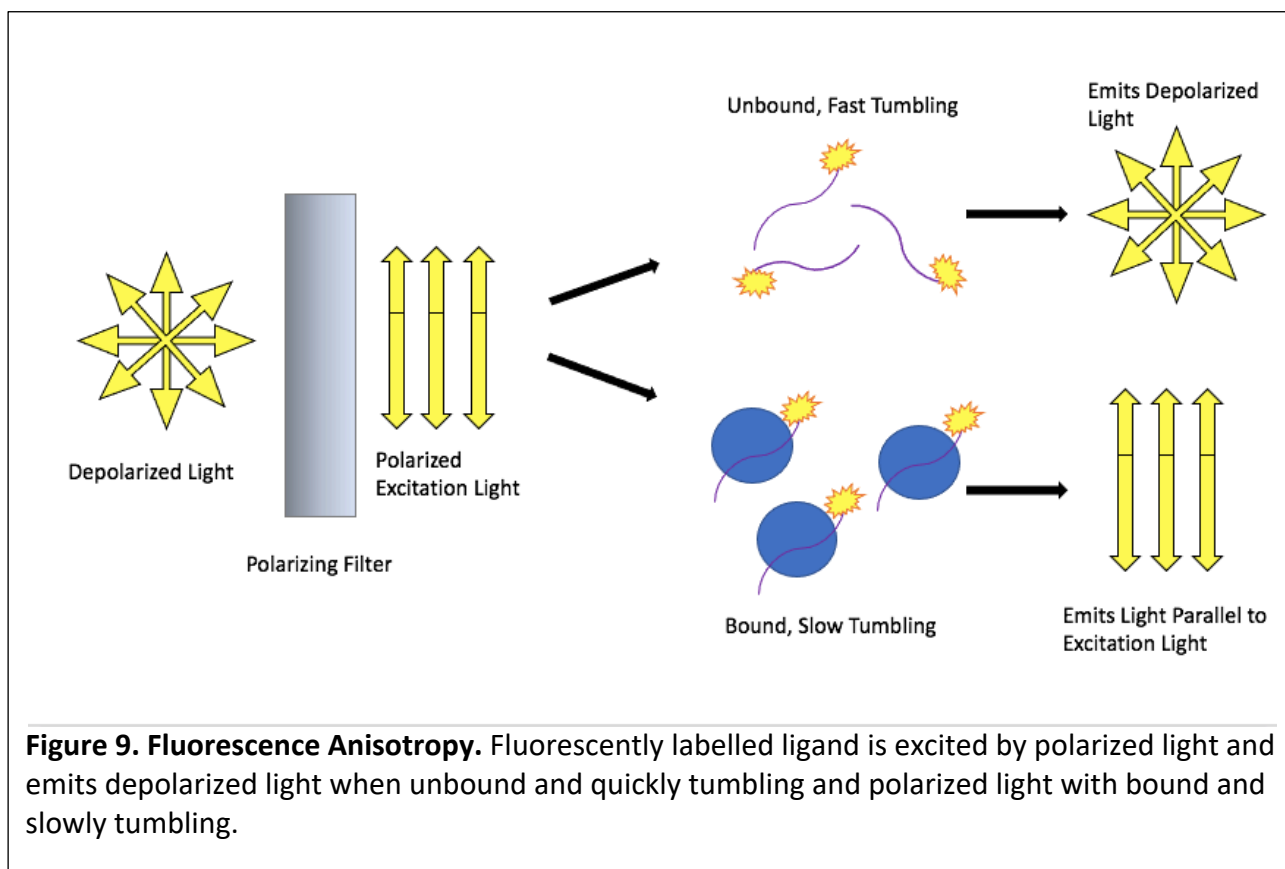


Figure 8. Urea gel with labelled RNA. Lane 1: Minus 1, 167 nt, Lane 2: Minus 2, 121 nt, Lane 3: Minus strand 3, 122 nt long, Lane 4: Minus 4, 82 nt, Lane 5: Minus 5, 131 nt

Binding measured by Fluorescence Anisotropy: I then tested CypA's affinity for the five minus sense strands in several Fluorescence Anisotropy (FP) assays. Fluorescence Anisotropy uses polarized light to excite the fluorophores attached to labelled ligands (in this case, the FTST on the RNA). At concentrations of protein below the K_d , most of the ligand is free and, after excitation, tumbles and emits light in different orientations. With higher protein concentration above the K_d , the ligand is bound in a larger complex so tumbles more slowly and emits a greater fraction of light parallel to the excitation light than in other orientations (see **Figure 9**).

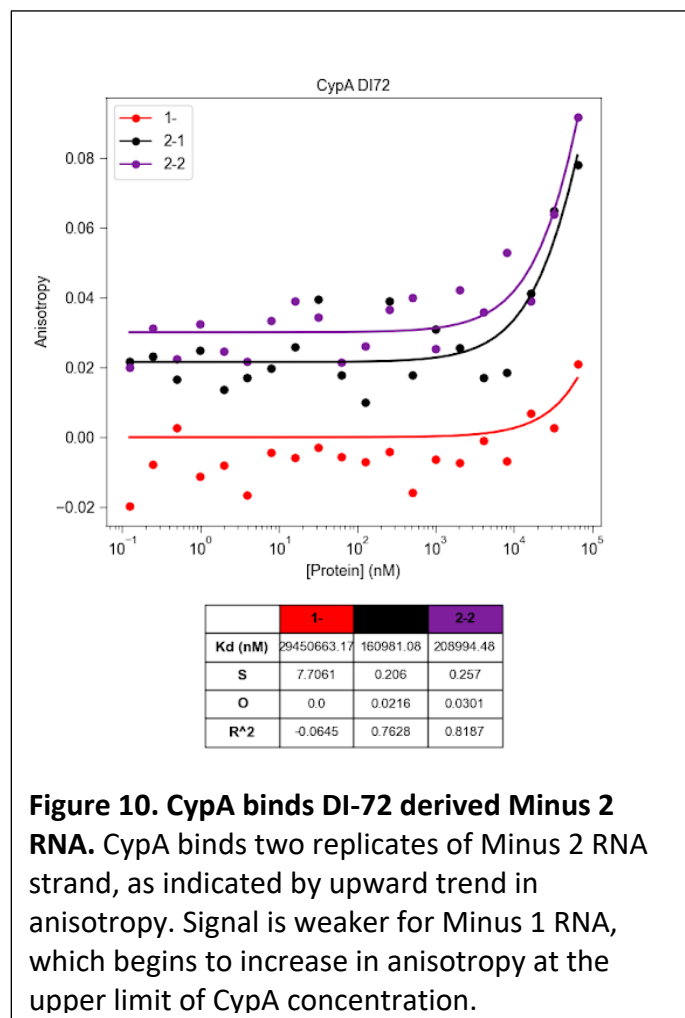
FP compares the intensity of perpendicular (free, tumbling) light to parallel (bound) light to calculate the fraction of ligand bound to the protein. Specifically, anisotropy is calculated as

$\frac{I_{par} - I_{per}}{I_{par} + 2 * I_{per}}$, where I_{par} is intensity of parallel light and I_{per} is intensity of perpendicular light.



The first two FA trials were run on all five RNA strands and a negative control (the biological RNA Gas5). These trials suggested an increased anisotropy as CypA concentration increased for the DI-72 derived RNA, but no shift in anisotropy for the negative control Gas5. These results also indicated preferential binding and implied binding specificity: CypA caused a shift in measured intensities for previously implicated RNA molecules, with different shifts for different sequences, but no shift for an arbitrarily selected negative control. However, the overall intensity of anisotropy for these trials decreased with CypA concentration (likely due to RNA crashing out of solution) and thus a low signal to noise ratio. These results were thus considered qualitative and the issue corrected in a final trial.

I conducted a final FA experiment with the two DI-72 RNAs that showed the most consistent anisotropic shifts between previous, qualitative trials: Minus 1 and Minus 2. **Figure 10** shows that while the Minus 1 strand shows an upward trend at the highest limit of CypA concentration, the Minus 2 strand undergoes a definite shift as CypA concentration is increased. Both the upper curves were generated with the same Minus 2 strand RNA at the same concentration and thus show similar offsets of background intensity. The bottom curve, showing binding to the Minus 1 strand, was



generated at a slightly lower concentration, and correspondingly has a lower offset of background intensity.

The trial tested two Minus 2 strand RNAs, which reassuringly produce anisotropy curves of similar trends. The K_d reported (about 185 μM averaged between Minus 2 strand curves) is a lower limit estimation, as the anisotropy curves do not plateau at the upper end of CypA concentration. This lower-level estimation suggests binding and is consistent with that reported by Kovalev et al.¹¹ (in the micromolar range). These initial FP assays are consistent with the literature and provide good evidence that CypA binds RNA *in vitro*. They also suggest some specificity, as CypA bound certain strands (Minus 2 being the best characterized) more tightly than others and did not bind the negative control Gas5.

In Vivo RNA Immunoprecipitation

Formaldehyde and UV RNA Immunoprecipitation

The first goal of my thesis was to verify, with a well-established equilibrium technique, that CypA is capable of RNA binding *in vitro*. The above results indicate such capabilities, as well as suggesting binding specificity. While ascertaining that CypA binds the DI-72 RNA is important and verified the RNA binding activity of CypA, a further goal of my thesis and the next pertinent problem was to identify what RNA molecules CypA binds in human cells, thus providing information as to the biological relevance of CypA-RNA interaction.

To determine the identity of RNA molecules that interact with CypA in human cells, I performed two types of RNA immunoprecipitations in HeLa cells: formaldehyde (f-RIP) and UV (UV-RIP). A strength of this approach is that CypA is expressed endogenously at high levels, precluding the use of strategies that require overexpression of the target for detection. This ensured that I was observing complexes present at natural conditions. These procedures used two different methods to crosslink (covalently link interacting molecules) cells before extracting the target protein. Formaldehyde (FA) crosslinking covalently links amine groups with high efficiency, allowing extensive linkage of molecules that are proximal but may not be directly interacting²⁵. UV cross-linking, which uses 254 nm light to irradiate cells, is more specific and cross-links directly interacting molecules (“zero distance linkage”). However, UV cross-linking efficiency is low: certain studies estimating only up to about 5% efficiency²⁶. Both formaldehyde and UV crosslinking immunoprecipitations were performed for deeper insight into CypA’s RNA interactome, as formaldehyde linkage is broad and more efficient and UV linkage is specific but less efficient.

The basic steps of an RNA immunoprecipitation are illustrated in **Figure 11**. Following crosslinking, lysed cells were incubated with a mouse derived, Immunoglobulin G (IgG) antibody specific to CypA (anti-CypA, ab58144). The antibody was previously verified by Western Blot. IgG antibodies are made up of four polypeptide chains: two heavy chains and two light chains that together form a flexible Y shape³¹. While the antigen binding region of the antibody is highly variable (to bind diverse antigens), the region of the two heavy chains at the bottom of the Y shape is highly conserved and can be targeted to precipitate antibody-antigen complexes³¹. After incubation with anti-CypA, magnetic beads specific to IgG antibodies were used to immunoprecipitate the antibody bound CypA that was covalently attached to RNA. A proteinase was then added to digest CypA, freeing RNA molecules, and RNA was purified from the samples (referred to as “pulldown” samples). An “input” sample of RNA was also taken and purified before addition of the antibody but after an initial incubation with beads. This

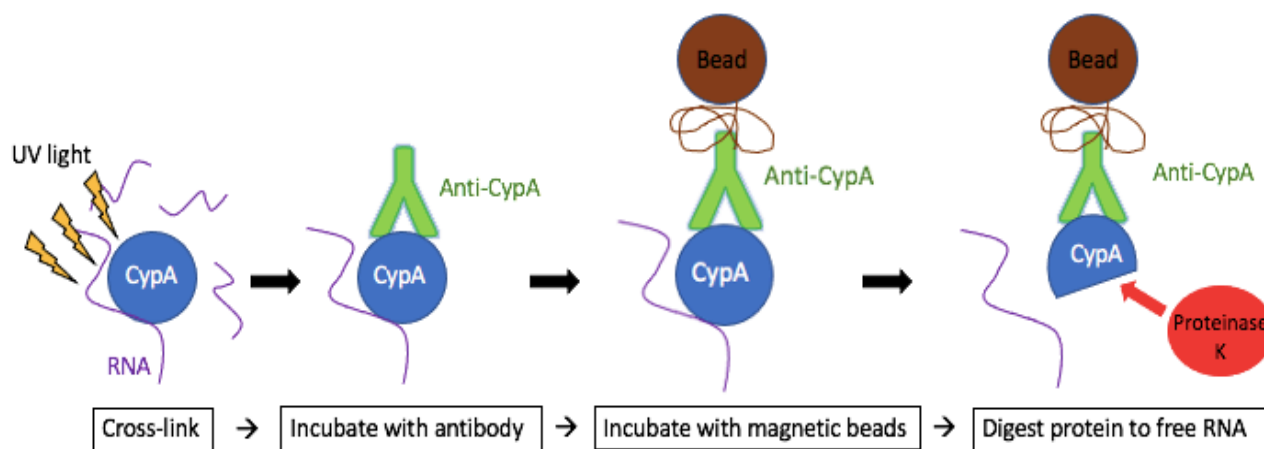


Figure 11. RNA UV Immunoprecipitation. Cells are crosslinked by irradiation with UV light, incubated with anti-CypA, incubated with magnetic beads that bind anti-CypA, pulled down, and RNA is detached by digestion of CypA and other proteins by Proteinase K.

initial incubation removed non-specific molecules that bound to magnetic beads and may thus have otherwise been pulled down with relevant RNA molecules. The result is a pool of RNA molecules that were pulled out due to a covalent interaction with CypA.

While some RNA-protein interaction purification techniques allow for sequencing of the binding site, the described approach frees the entire interacting RNA molecule. Thus, after library preparation and sequencing, only information about the full length RNA molecule is available, not the binding site.

RNA Sequencing

RNA libraries for fRIP and UV-RIP experiments were prepared for sequencing by fragmentation to desirable read size (~300 nt) using heat and magnesium ions (Mg^{2+}). RNAs have catalytic activity and thus the capacity for self-cleavage^C. Heat and metal ions accelerate the process of self-degradation and Mg^{2+} in particular has been shown to form complexes with RNA molecules to stabilize certain structures which are conducive to RNA catalytic cleavage^C. Library preparation continued after self-fragmentation of RNA with reverse transcription (converted molecules to complementary DNA), ligation of double stranded DNA Illumina adapters, and PCR (polymerase chain reaction) amplification. Illumina adapters contain sequences necessary for 1) adhesion to an Illumina sequencing flow cell by complementary base pairing and 2) “barcoding” or labelling molecules to identify what sample they came from. This library prep also included the addition of UMIs (Unique Molecular Identifiers), which are unique sequences of nucleotides given to each molecule. These are useful because of the error inherent in amplifying libraries for sequencing—certain molecules may be preferentially

amplified by PCR and thus make up a larger portion of sequencing reads. By counting UMIs rather than sequences, it is possible to accurately determine how many times an identical RNA sequence occurred in each sample. An illustration below (**Figure 12**) clarifies the use of UMIs.

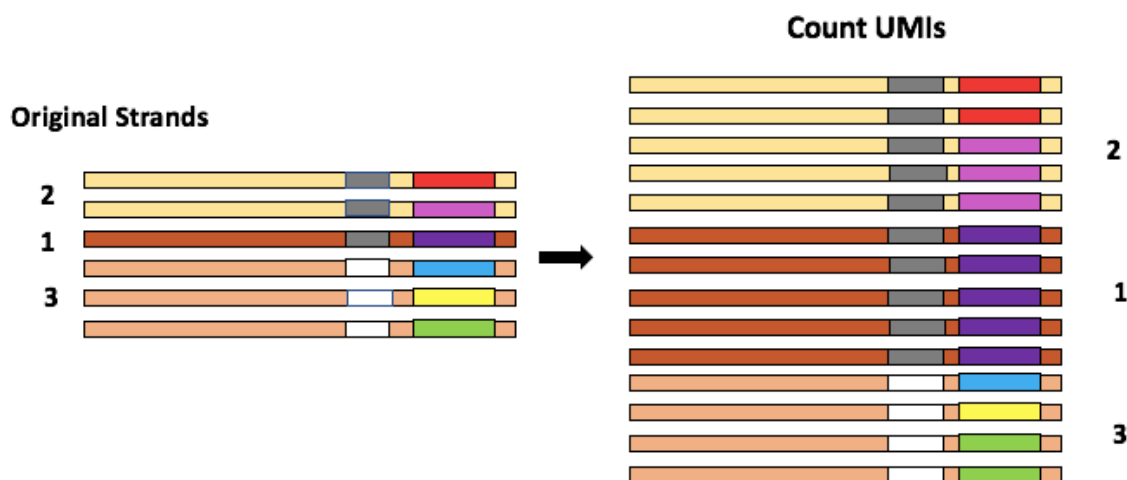


Figure 12. After amplification, UMIs allow accurate counts. Each different color represents a different RNA sequence. All unique RNA molecules (same sequence or not) will be tagged with a unique sequence of nucleic acids. After duplication, the number of UMI sequences is the number of original strands of any sequence the sample contained. Each RNA molecule also contains an adapter (grey or white) that indicates what sample it was from.

Six samples of RNA were sequenced: an input and pulldown from two FA experiments and one UV experiment. Amount of reads per sample ranged from 22,917,431 to 76,393,313 and 86% of reads across samples had a quality score > Q30. Quality scores are calculated based on the number of unidentifiable nucleotides in sequences: a quality score of Q30 represent an average error of 1 in 1,000 nucleotides. Both UV and FA inputs and pulldowns (pre- and post-immunoprecipitation) were processed by removing low-quality and duplicate reads (using UMIs) and aligning unique reads to the human genome (version GRCh38). Unique reads per genomic feature were then counted.

Determination of Statistically Significant Enrichment and Depletion

I then determined which genes were enriched in pulldown samples as compared to input samples using DESeq2²¹. DESeq2 normalizes counts per gene based on total read counts across all genes for a particular sample and averages the normalized counts per gene across sample replicates, which accounts for library size differences. It then calculates a logarithmically scaled fold change: $\log_2(\text{average normalized pulldown} / \text{averaged normalized input})^{21}$. For each gene count, DESeq also calculates p-values for enrichment using the Wald test with the null hypothesis that there was no differential expression across sample groups (pulldown and input)²¹. P values were adjusted using the Benjamini and Hochberg method to scale p-values and reduce the number of false positives given a certain p-value cutoff²¹. As it uses averaged read counts, DESeq2 requires replicate experiments. I was able to conduct two formaldehyde immunoprecipitations but only one for UV, so consider just the FA enriched genes in the following analysis.

I considered as significant genes with an adjusted p-value of less than 0.05. **Figure 13** shows a plot with all genes comparing their significance to their log2 fold change. The negative of the \log_{10} of adjusted p-values is plotted on the y-axis such that the higher the gene, the more significant its fold change. The outward and upward sweep of this plot from the origin reveals a general trend for enriched genes: the greater magnitude of the fold change, the more statistically significant the calculation of fold change. This increases confidence in the likelihood

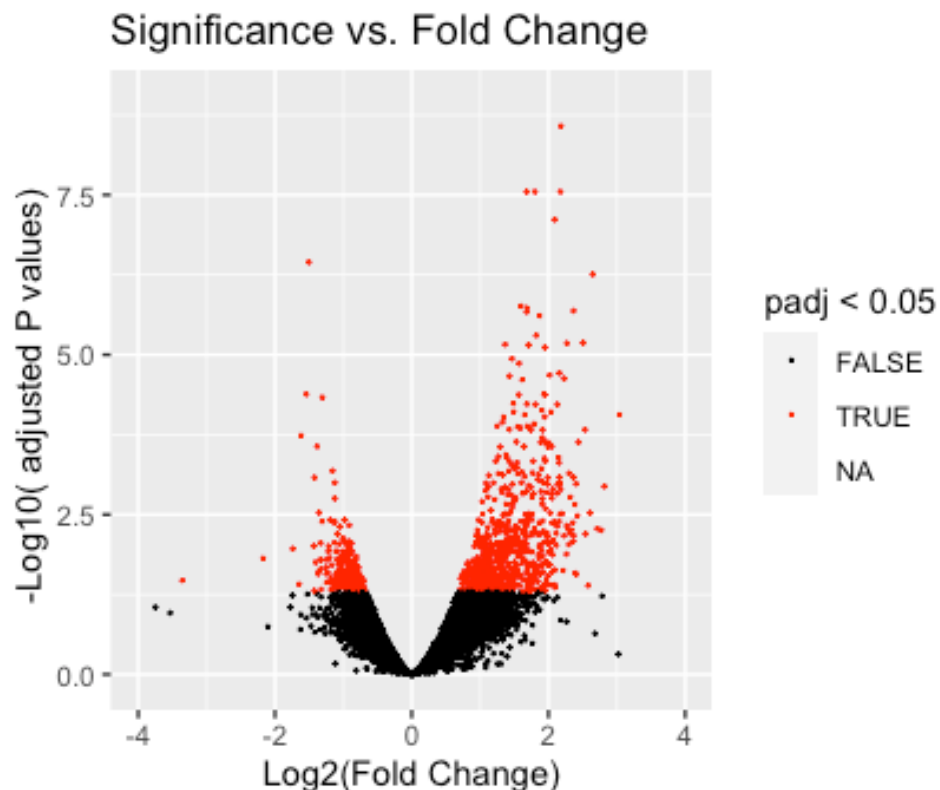


Figure 13. Significance vs. Fold Change of all Genes. Log₂ of Fold Change is plotted on the x-axis with $-\log_{10}$ of adjusted p-value on the y-axis for easy visualization of statistically significant genes. Genes with adjusted p-values of less than 0.05 are plotted in red. Genes to the right of the origin (positive log₂ fold change) are enriched, while genes to the left are depleted in pulldown vs. input.

that transcribed genes with a higher fold change are indeed interacting with CypA. Figure 8, then, indicates that some transcribed genes are significantly enriched (positive log₂ fold change) and others significantly depleted (negative log₂ fold change) with respect to CypA interaction, as expected.

Clustering of Enriched Genes

There were 709 enriched genes observed to have a positive log₂ fold change and an adjusted p-value of less than 0.05. While this large number provides little information as an unannotated list of genes, patterns which emerge after grouping these genes by biological function annotation reveal specific, enriched pathways. These pathways may suggest a functional role for the RNA-binding activity of CypA.

The DAVID (Database for Annotation, Visualization and Integrated Discovery) clustering tool was used to identify several biological functional groups in which enriched genes occurred²². DAVID calculates similarity between genes in a submitted list by comparing gene annotation terms (such as biological function and pathway) obtained from its extensive annotation database²². It then clusters genes based on similarity distances²². DAVID clustered the 709 enriched genes into groups based on biological function and calculated the significance for each group (based on how many genes belonging to a functional group occurred in the given gene set compared to how many genes would occur in that group by chance). **Figure 14** shows the top clustering results with bar height corresponding to the number of genes that occurred in each group and color to adjusted p-value. Note that this p-value is associated with the significance of the functional group, not the enrichment of the genes within it.

CypA interacts with genes involved in transcription, transcriptional regulation, and kinase activity with the highest significance. There is overlap between top functional groups (zinc fingers and plexins are metal binders, zinc fingers are transcription factors, ATP-binders can be kinases). The diversity of cellular location in functional groups, with cell-to-cell adhesion (extra-cellular) as well as transcriptional regulation (nucleus) is unsurprising due to the ubiquity of CypA: it is found in the nucleus, cytoplasm, and is secreted⁷. In the nucleus, CypA is known to mediate translation through interactions with major transcription factors, specifically YY1⁸ and NF- κ B.⁹ It has also plays a role in kinase signaling by proline-directed regulation of such molecules as the tyrosine kinase Itk¹⁹. That its most significant set of interactions is with genes coding for proteins involved in transcription, transcriptional regulation, and kinase activity is therefore consistent with our knowledge of the functional roles of CypA.

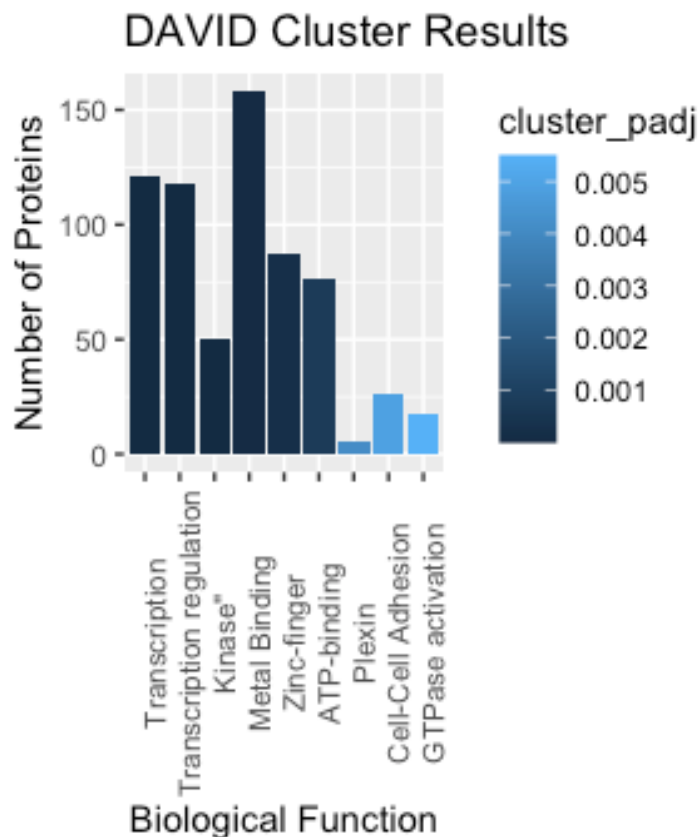


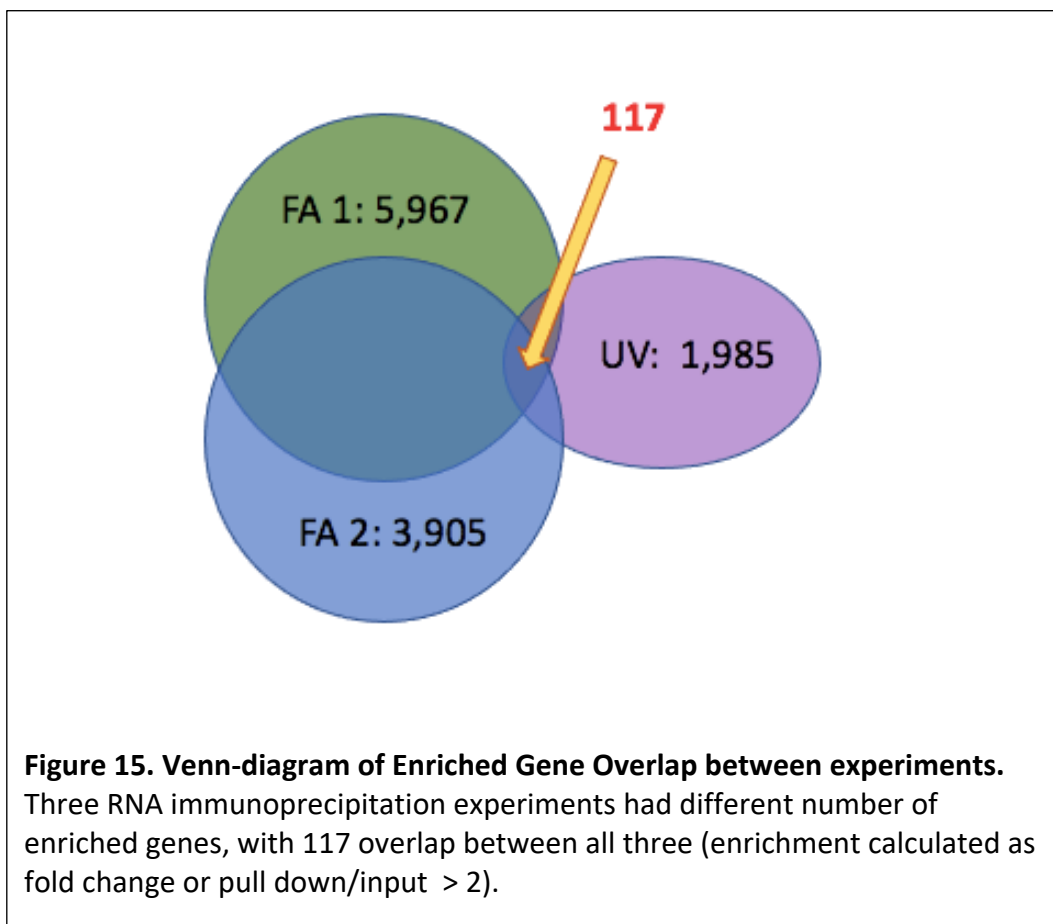
Figure 14. DAVID Clustering Results. Top significant biological functional clusters for the enriched DESeq2 genes were identified with DAVID. Bar heights are number of genes belonging to each cluster, colored according to adjusted p-value for that cluster.

One important caveat should be considered for this set of formaldehyde-linked interactions: formaldehyde links broadly and some of the enriched genes may be due to indirect interactions, or RNA molecules that a protein partner of CypA interacts with. NF- κ B, for example, regulates immune system response genes²² and Rab GTPases are known to influence inflammation and immunity²⁷. Among the set of GTPases, then, could be indirect interactors of CypA. YY1, another known CypA mediated protein⁸, forms homodimers that are “stabilized by low specificity RNA binding.”²⁸ The diverse set of enriched formaldehyde genes may thus contain YY1 and NF- κ B RNA interactions, as well as other indirect or secondary interactions. Thus, in the future it will be critical to cross-reference the formaldehyde crosslinking data with

the UV crosslinking data to determine a refined set of RNA targets that are due to direct interactions.

Consideration of Robustly Enriched Genes

The above analysis does not take into consideration the UV-crosslinked sample, as only one replicate was available. To take advantage of the rich insight that UV-crosslinking would provide, I ran my own analysis to calculate enrichment. I first normalized read counts for each gene by dividing gene counts by “counts per million,” or total number of counts for each gene’s sample divided by one million. I then calculated fold change (pulldown/input) for each gene (treating FA samples and UV samples separately). I considered as enriched genes with a fold



change greater than 2. There were 5,967 such enriched genes for one FA trial, 3,905 in another, and 1,985 in the UV trial (**Figure 15**). 117 genes appeared with a fold change of greater than 2 in all three samples (FA1, FA2, UV).

Comparison of these results with enrichment data from DESeq2 analysis yielded 13 genes that were consistently enriched across methods of analysis and samples, referred to as robustly enriched genes (REGs). Other notable genes that emerged were seven from DESeq2 analysis with a strikingly low adjusted p-value ($p_{adj} < 10^{-6}$), referred to as strikingly significant genes (SSGs). Next to determine was how the functionality of these two exceptional gene sets relate to known CypA activity. **Figure 16** shows the groups plotted in red (SSGs, extremely low p_{adj} in FA samples) and green (genes enriched in FA and UV samples) over other DESeq2 enriched genes (positive fold change, adjusted p-value < 0.05).

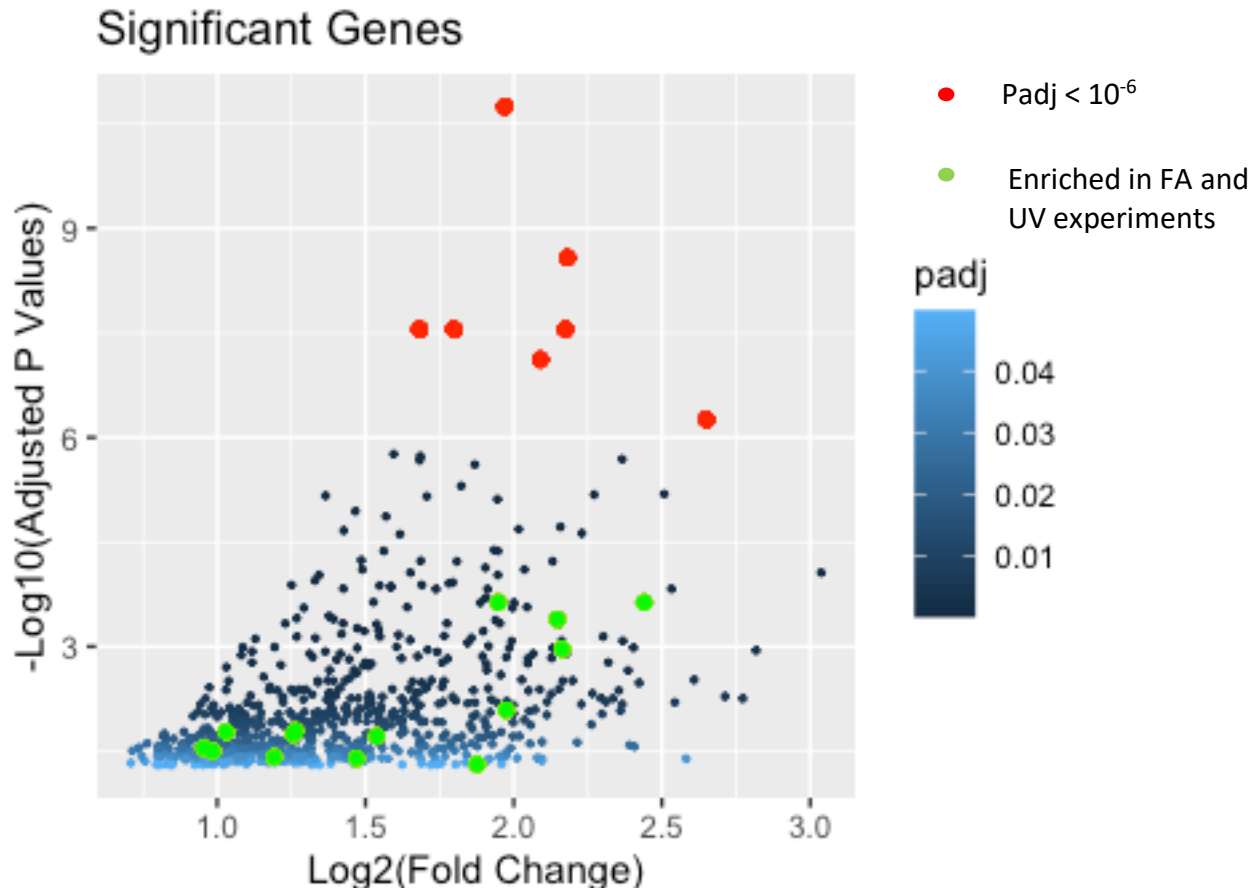


Figure 16. FA enriched genes with overlaid prominent significant genes. 709 significantly enriched genes from DESeq2 analysis (positive log₂ fold change and p adj < 0.05) are plotted, colored by adjusted p value. Highlighted are genes that occurred in all methods of enrichment calculation (REGs, green) or had extremely low adj p-values (SSGs, red).

Of the 20 genes belonging to either of the two enriched gene sets, there were 15 with biological roles relating to key CypA activity, including proto-oncogenes and transcription factors. Figure 17 (with information from the UniProt database²⁹) summarizes the functions of the SSGs and REGs. Overexpression of CypA has been shown to stimulate cancer cell growth in various studies by enabling cancer proliferation, blocking apoptosis, and regulating cell cycle progression⁷. Four of the SSGs and five of the REGs have been implicated in cancer or abnormal

cell growth, including *SKI*, which codes for a nuclear proto-oncogene²⁹, *MEGF6*, which induces metastasis²⁹, and *RXRA*, a nuclear receptor that has been implicated in renal cancer²⁹. CypA also plays a critical role in the transcriptional activity of several key transcription factors. It has been shown to interact specifically with the previously mentioned transcription factor YY1, thereby altering YY1's activity⁸. Also mentioned, it has been shown to interact with a subunit of the general transcription factor NF- κ B⁹, which regulates genes involved in immune system response²². Three SSGs and four of the REGs are involved in transcription and gene regulation.

Figure 17. SSGs, REGs and their functions. Functional Information obtained from UniProt²⁹

Gene Name	Enrichment Set	Function	Implicated in Cancer/Abnormal Cell Growth?	Implicated in Transcription/Gene Regulation?
<i>SKI</i>	SSG	Nuclear proto-oncogene	X	
<i>MEGF6</i>	SSG	epidermal growth factor, induces metastasis	X	
<i>RXRA</i>	SSG	Nuclear receptor, Host-virus interaction, Transcription, implicated in renal cancer	X	X
<i>CAPN15</i>	SSG	Zinc-finger repeats, transcription factor		X
<i>ZNF469</i>	SSG	Zinc-finger, may be transcription factor		X
<i>MIRLET7BHG</i>	SSG	Long non-coding RNA, related pathways include metastatic brain tumor	X	
<i>SETD1A</i>	REG	Histone Lysine Methyltransferase component, marks TSS of active genes		X

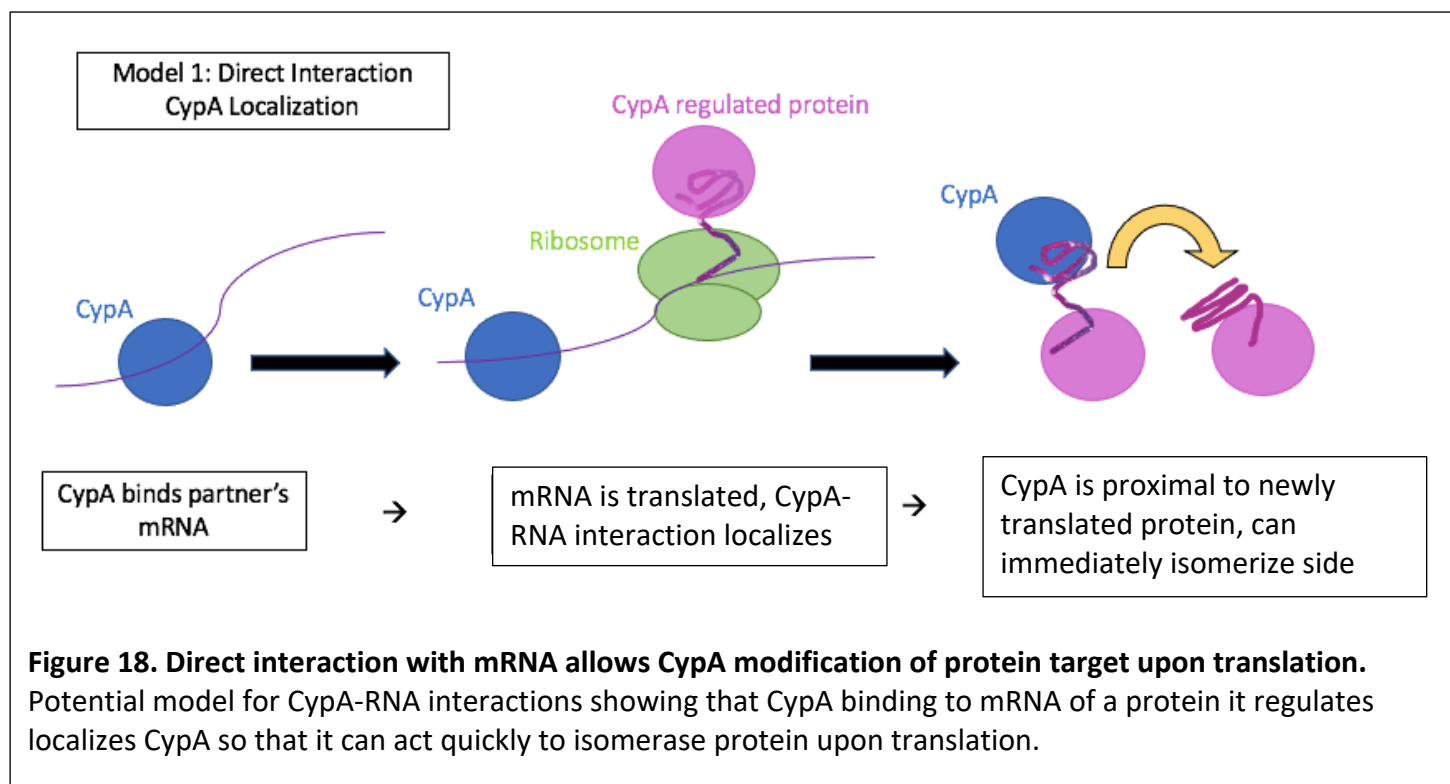
<i>PHRF1</i>	REG	Contains PHD and ring-finger, chromatin mediated gene regulation		X
<i>GAS8</i>	REG	Binds tyrosine-protein kinase receptors, which are implicated in cell growth and migration, target in cancer therapy	X	
<i>CBX4</i>	REG	SUMO-protein ligase, indirectly regulates p53/TP53 transcriptional activation		X
<i>MUC1</i>	REG	Cell-surface glycoprotein, overexpressed in many cancers	X	
<i>GTF2F1</i>	REG	General transcription initiation factor, binds to RNA polymerase II		X
<i>RABL6</i>	REG	GTPase, may play a role in cell growth, oncogene	X	
<i>GADD45B</i>	REG	Expression increased with growth arrest and DNA damage	X	
<i>BRD3</i>	REG	Contains bromodomain, recognizes acetylated lysine residues of histones (chromatin mediated gene regulation)	X	
<i>SF3A2</i>	REG	pre-mRNA splicing, component of splicing factor SF3A		
<i>AP3D1</i>	REG	facilitates vesicles budding from the golgi membrane, and may be directly involved in		

		trafficking to lysosomes		
<i>BAIAP2</i>	SSG	Adaptor protein, links membrane G-proteins to cytosolic		
<i>AC011498.7</i>	REG	unknown		
<i>C13orf46</i>	REG	unknown		

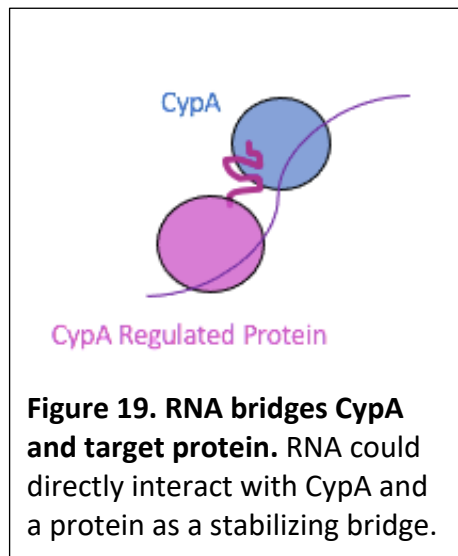
The involvement of SSGs and REGs in cancer and cell growth and/or transcription and gene regulation is striking as CypA plays a major role in these two areas: it is plausible that CypA indeed regulates these genes by interacting with their RNA transcripts.

Potential Models for CypA-RNA Interaction

The presented results are particularly exciting in showing that genes enriched in pulldown samples are involved in CypA-regulated pathways. As CypA has been shown to mediate the transcriptional activities of YY1⁸ and NF- κ B⁹ as well regulate the tyrosine kinase Itk,¹⁹ it may similarly interact with other transcription factors and kinases. An important distinction must be made, however, between the mRNA transcripts that encode these proteins and the proteins themselves: the immunoprecipitation experiments determined enrichment for RNA, not protein, levels. An intriguing mechanism by which CypA could interact with both the mRNA and protein product of a certain gene is localization by direct RNA-interaction (Figure 18). CypA could potentially bind the mRNA of the protein it regulates, thus localizing to the area of translation, and be proximally available to edit the side-chain orientation upon (or during) translation. This would accelerate synthesis of fully folded proteins.

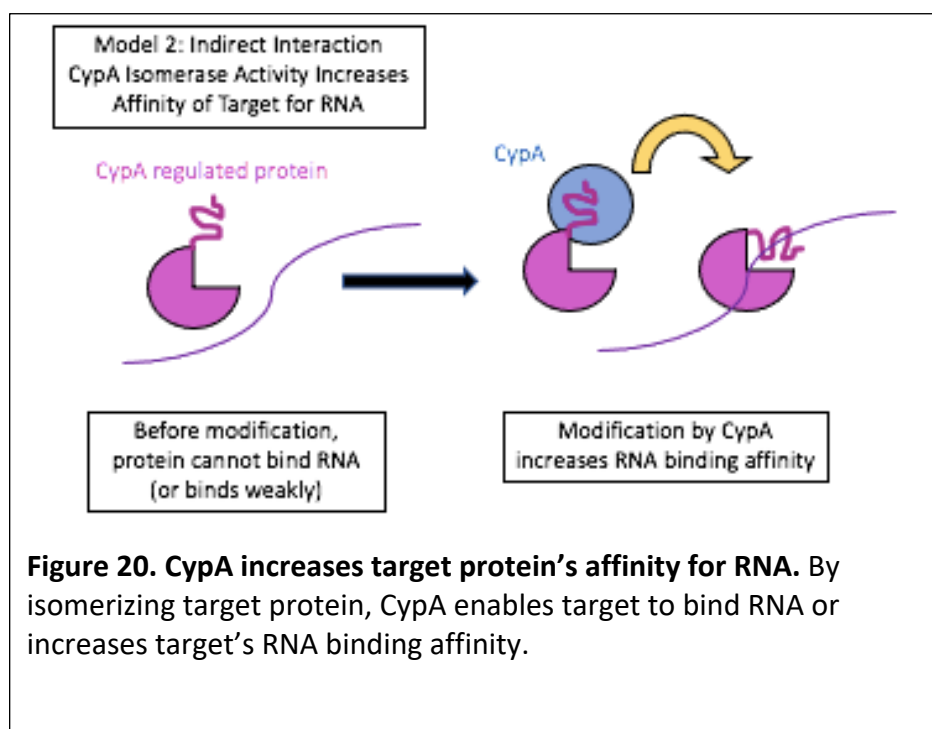


RNA could also act as a bridge between CypA and a protein that it regulates, increasing stability of the complex and efficiency of isomerization. This would be consistent with previous experiments in the lab that showed random RNAs added to an activity assay increased CypA's rate of isomerization¹⁰: in bringing proteins together and stabilizing their positions, RNA could increase the rate at which CypA finds and acts on the proteins it regulates (see **Figure 19**).



Perhaps, though, it is not direct interaction but rather indirect mediation of protein-RNA binding that implicates CypA in certain gene pathways. A study has shown that interaction between a catalytically active CypA and NS5A, a protein necessary for hepatitis C viral replication, stimulates NS5A's RNA binding¹⁷. Perhaps, then, CypA's isomerase activity changes the shape of certain proteins to increase their affinity for or enable RNA binding. In this case,

the above FA pulldown experiments would include many indirect interactions. **Figure 20** illustrates the potentiality of indirect CypA-RNA interactions.



Future Directions

The presented work merely scratches the surface of CypA-RNA interactions, leaving many interesting questions to be addressed. A first step would be to identify certain RNA sequences and/or structures that contribute to the observed specificity of CypA binding. To do this, a similar immunoprecipitation could be performed with the additional step of RNA digestion before sequencing and protein digestion: if RNase (protein that digests RNA) is added to the sample before RNA is freed from CypA binding, the RNA sequestered by protein binding will be protected from digestion, leading to sequencing of merely the RNA binding site. Then, these sequences could be tested *in vitro* to verify CypA affinity. Further, *in vitro* activity assays could be done with an RNA sequence that binds with reasonable affinity to ascertain RNA's effect on CypA isomerase activity.

To investigate if isomerase activity of CypA is required *in vivo* interactions (indirect or direct), an RNA immunoprecipitation could be done with catalytically dead CypA. The CRISPR-Cas9 system could be used to introduce a mutation removing catalytic abilities into the CypA gene of HeLa cells, pulldown performed, and results compared with those reported above. As CypA is an extremely important cellular protein, however, the viability of the cells may be impacted by removing its isomerase activity. An alternative method would transfect a vector with a catalytical dead CypA mutant and an attached his-SUMO tag into HeLa cells. Then, a his-SUMO specific probe rather than an antibody could be used for pulldown. Comparing RNAs pulled down with a catalytically dead CypA to those pulled down with a catalytically active CypA would provide information on the RNA binding that is related to or depends on CypA's

isomerase activity. This would provide evidence for a previously unknown level of RNA-protein regulation.

Conclusion

RNA-protein interactions have numerous and varied critical biological roles, many yet undiscovered.^{1,2,3,4} Here, the novel CypA-RNA interaction have been broadly explored, providing *in vitro* and *in vivo* evidence for CypA-RNA binding. Fluorescence Anisotropy assays with segments of viral DI-72 RNA derived from the genome of the TBSV suggest CypA-RNA binding specificity *in vitro*, as the negative control RNA did not bind CypA while several Minus Strand DI-72 transcripts showed anisotropic shifts indicative of binding. Specific binding, perhaps relating to the sequence or structure of the RNA molecule, is exciting as it could be evidence that RNA plays a functional role in CypA activity. Several RNA immunoprecipitations using both UV and formaldehyde cross-linking methods produced a set of statistically enriched genes in human cells, the most significant being transcription related genes. A select group of enriched genes which appeared in all formaldehyde and UV RNA immunoprecipitation samples or had extremely low adjusted p-values are especially interesting as their proposed functions correspond to key CypA activities: transcriptional regulation and cancer progression. While these initial studies show that CypA binds different RNAs with different affinity, further work remains to identify specific RNA characteristics (sequences or structures) that contribute to CypA's preferential binding and determine reliable K_d s for these interactions. Next, activity assays and RNA immunoprecipitations with catalytically dead CypA will increase our understanding of the role RNA plays in CypA activity. This initial exploratory study adds to the body of research implicating CypA as an unexpected yet important RNA binder and opens exciting avenues for future research on CypA-RNA interactions and other novel RNA binding proteins.

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