ARiBo pull-down for riboproteomic studies based on label-free quantitative mass spectrometry

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ABSTRACT

As part of their normal life cycle, most RNA molecules associate with several proteins that direct their fate and regulate their function. Here, we describe a novel method for identifying proteins that associate with a target RNA. The procedure is based on the ARiBo method for affinity purification of RNA, which was originally developed to quickly purify RNA with high yields and purity under native conditions. The ARiBo method was further optimized using in vitro transcribed RNA to capture RNA-associating proteins from cellular extracts with high yields and low background protein contamination. For these RNA pull-downs, stem–loops present in the immature forms of let-7 miRNAs (miRNA stem–loops) were used as the target RNAs. Label-free quantitative mass spectrometry analysis allowed for the reliable identification of proteins that are specific to the stem–loops present in the immature forms of two miRNAs, let-7a-1 and let-7g. Several proteins known to bind immature forms of these let-7 miRNAs were identified, but with an improved coverage compared to previous studies. In addition, several novel proteins were identified that better define the protein interactome of the let-7 miRNA stem–loops and further link let-7 biogenesis to important biological processes such as development and tumorigenesis. Thus, combining the ARiBo pull-down method with label-free quantitative mass spectrometry provides an effective proteomic approach for identification of proteins that associate with a target RNA.

Keywords: RNA pull-down; proteomics; RNA–protein interactions; pre-miRNA; let-7 biogenesis

INTRODUCTION

RNA molecules associate with proteins to regulate many biological processes, and the identification and characterization of ribonucleoprotein complexes is currently an area of intense scientific interest. For profiling interaction partners of macromolecules of interest, affinity purification coupled to mass spectrometry (AP-MS) has become a standard method. Until recently, the majority of efforts for characterizing interactions of RNA–protein networks by AP-MS have focused on isolating complexes using affinity-tagged proteins (Oeffinger 2012; Faoro and Ataide 2014; McHugh et al. 2014). However, there is also a need to characterize RNA–protein networks from the vantage point of the RNA, and a few strategies have been developed to purify RNA–protein complexes using affinity-tagged RNAs. Although most of these AP-MS investigations using tagged RNAs are based on in vitro assembly with proteins from cellular extracts, the characterization of RNA–protein complexes formed in vivo has also been investigated (Hogg and Collins 2007a,b; Vasudevan and Steitz 2007; Said et al. 2009; Baltz et al. 2012; Castello et al. 2012; Oeffinger 2012; Kwon et al. 2013; Faoro and Ataide 2014; Gersterber et al. 2014; Kotzer-Nevo et al. 2014; Leppke and Stoecklin 2014; McHugh et al. 2014; Beckmann et al. 2015; Dong et al. 2015). Despite these efforts, there is still much that remains to be learned about the protein-interaction networks of specific RNAs.

Quantitative approaches in AP-MS studies are very powerful for the comprehensive and reliable identification of molecular interaction partners. Although the SILAC (stable isotope labeling by amino acids in cell culture) metabolic labeling approach has contributed significantly to the identification of interaction partners for specific proteins and RNAs as part of MS-based quantitative proteomics studies (Chen et al. 2015), label-free methods are becoming increasingly popular because they are usually simpler and cheaper (Tate et al. 2013; Sandin et al. 2014). Moreover, the recent development of high-resolution liquid chromatography coupled with tandem mass spectrometry methods (LC–MS/MS) allows label-free quantification with high accuracy and reproducibility (Skarra et al. 2011; Choi et al. 2012; Cox et al. 2014; Teo et al. 2014). When coupled with robust quantitative
methods for data analysis, label-free AP-MS methods allow for the possibility to computationally remove nonspecific background interactions using a negative control, such as an RNA pull-down performed with a variant form of the target RNA.

Several affinity tags can be attached to an RNA of interest for its purification with associated proteins, and they include biotin, fluorophores, RNA aptamers that bind small molecules, and RNA recognition elements of proteins or protein domains. The RNA-based tags are particularly attractive for AP-MS applications given their suitability for purification under native conditions in both in vitro and in vivo applications (Sirisawat and Engelke 2002; Oeffinger 2012; Faoro and Ataide 2014; McHugh et al. 2014; Panchapakesan et al. 2013). Commonly used RNA-based tags for AP-MS studies include streptavidin and tobramycin aptamers (Hartmut et al. 2002; Hogg and Collins 2007b; Nelson et al. 2007; Butter et al. 2009; Leppke and Stoecklin 2014), as well as pp7 and MS2 RNA hairpins (Hogg and Collins 2007b; Nelson et al. 2007; Said et al. 2009; Slobodin and Gerst 2010). Some of these tags have been combined for tandem affinity purification in order to increase the purity of the isolated macromolecular complexes and reduce nonspecific interactions (Hogg and Collins 2007b; Nelson et al. 2007). However, it has become clear that these multistep affinity purification protocols may lead to dissociation of weak and/or transient interactors, which would significantly reduce the number of interactors identified. Thus, there is considerable interest in developing quick single-step affinity purification protocols that maximize the capture of true interaction partners and minimize nonspecific interactions (Oeffinger 2012; Faoro and Ataide 2014; McHugh et al. 2014). Moreover, given that only a few studies have combined RNA pull-down with quantitative MS (Butter et al. 2009; Lee et al. 2013; Adachi et al. 2014; Leppke and Stoecklin 2014), there is a need to develop robust RNA affinity purification procedures that can complement recent developments in label-free quantitative proteomics (Oeffinger 2012; Faoro and Ataide 2014; McHugh et al. 2014).

Over the past few years, our laboratory has optimized the ARiBo procedure for affinity purification of in vitro synthesized RNAs with a 3′-ARiBo tag, which contains an Activatable glmS Ribozyme and the BoxB RNA from bacteriophage λ (Di Tomasso et al. 2011, 2012a,b). The λboxB RNA part allows the specific immobilization of an ARiBo-tagged RNA on Glutathione-Sepharose (GSH-Sepharose) resin via its high affinity to the λN-glutathione-S-transferase (GST) fusion protein, whereas the glmS ribozyme part can be activated by glucosamine-6-phosphate (GlcN6P) to liberate the RNA of interest and concomitantly produce a homogeneous 3′-end. Importantly, our ARiBo procedure rapidly generates highly pure RNA with very good yields under native conditions. Moreover, we have demonstrated that this procedure can be used to purify RNA with different sequences, secondary structures and sizes. In addition, it can be combined with complementary approaches to ensure 5′-homogeneity of the purified RNA (Salvail-Lacoste et al. 2013; Di Tomasso et al. 2014). Thus, the ARiBo procedure represents an attractive method for the purification of RNA–protein complexes in RNA-based AP-MS studies.

In this manuscript, we have optimized the ARiBo affinity purification method for riboproteomic studies based on label-free quantitative mass spectrometry. The RNA pull-down procedure was developed using in vitro transcribed ARiBo-tagged stem–loops present in the immature forms of miRNAs (miRNA stem–loops) to capture RNA-associating proteins from whole cell extracts (WCEs). Stem–loops derived from the precursors of let-7a-1 and let-7g were used (Bussing et al. 2008; Roush and Slack 2008; Viswanathan and Daley 2010; Thornton and Gregory 2012; Nguyen and Zhu 2015; Rehfeld et al. 2015), since several proteomic studies have been reported for these RNAs (Heo et al. 2008, 2009; Michlewski et al. 2008; Viswanathan et al. 2008; Michlewski and Caceres 2010; Chang et al. 2013; Lee et al. 2013). In addition, let-7a-1 and let-7g are two of the 12 human let-7 miRNAs that play important roles in mammalian development, metabolism, and cancer (Bussing et al. 2008; Roush and Slack 2008; Viswanathan and Daley 2010; Thornton and Gregory 2012; Nguyen and Zhu 2015; Rehfeld et al. 2015), and there is still significant interest in identifying proteins that control biogenesis of these miRNAs through interactions with the stem–loop structures present in their immature forms. We performed quantitative LC–MS/MS of RNA pull-downs using biological triplicates and two experimental controls to identify proteins that specifically bind to the stem–loops of let-7a-1 and let-7g. Several proteins were identified that were previously shown to bind immature forms of let-7 miRNAs (Heo et al. 2008, 2009; Michlewski et al. 2008; Viswanathan et al. 2008; Michlewski and Caceres 2010; Chang et al. 2013; Lee et al. 2013). In addition, we identified an extensive group of novel protein factors not previously found to bind these RNAs. Taken together, our results make an important contribution to defining the protein interactome of let-7 miRNA stem–loops. In addition, they demonstrate that combining the ARiBo pull-down with label-free quantitative MS represents a powerful approach for the identification of proteins that associate with a target RNA.

RESULTS

Optimization of the RNA pull-down assay

The ARiBo procedure for affinity purification of RNA was adapted to isolate proteins from cell extracts that specifically associate with a target RNA (Fig. 1). The initial target RNA that we tested was SL-let-7g, the stem–loop structure found in the immature forms of the let-7g miRNA (Fig. 2). The SL-let-7g RNA was first synthesized by in vitro transcription with an ARiBo tag at its 3′-end (Di Tomasso et al. 2011). The ARiBo-tagged SL-let-7g was then bound to the λN-GST
to the ARiBo-tagged SL-let-7g, whereas much lower levels of proteins were eluted in high abundance that specifically bound ARiBo-tagged SL-let-7g to that of a mock purification performed in the absence of a cell extract (Fig. 3B). To evaluate the specificity of the RNA pull-downs, we compared the protein elution profile from the purification with the ARiBo-tagged SL-let-7g to that of a mock purification performed with the ARiBo tag only. Remarkably, several proteins were eluted in high abundance that specifically bound to the ARiBo-tagged SL-let-7g, whereas much lower levels of proteins were eluted from the mock purification (Fig. 4A). Given that let-7 stem–loops are known to directly bind to the Lin28A protein (Piskounova et al. 2008; Desjardins et al. 2011, 2014), we used Western blotting with an anti-Lin28A antibody to assess the specificity of the RNA pull-down (Fig. 4B). We confirmed by Western blot analysis of several biological replicates that the Lin28A protein specifically associates with the ARiBo-tagged SL-let-7g (Fig. 4C) in comparison with the ARiBo tag control. Given that the Lin28A protein binds with high affinity to pre-let-7g (Kd of 0.15 nM) but also binds with lower affinity and specificity to other RNA sequences (Piskounova et al. 2008; Desjardins et al. 2011, 2014), the low levels of Lin28A in the mock pull-downs relative to the SL-let-7g pull-downs validate the specificity of the procedure.

The conditions of the RNA pull-down assays reported in Figure 4 were optimized to maximize the total amount of proteins eluted from the target RNA sample while reducing the total amount of proteins eluted from the ARiBo tag control (mock; Fig. 4A). Thus, these conditions contribute to discriminate true protein interactors from nonspecific background contaminants. We determined that salt concentrations of 5–10 mM MgCl2 and 100–250 mM NaCl are optimal in the load and wash buffers. In addition, we found that following incubation with the cell extract at least one wash step is necessary to reduce nonspecific binding. Subsequent wash steps are not desirable as they significantly reduced the total amount of protein in the RNA pull-downs,

**FIGURE 1.** ARiBo pull-down method for the affinity capture of RNA–protein complexes. (A) General scheme for isolating RNA-associating proteins from a WCE by RNA pull-down with an ARiBo tag. (B) Schematics of the ARiBo-tagged RNA showing the sequence and secondary structure of the RNA (gray shading) in its variable P1 stem and stabilize the P2 stem. The black arrowhead identifies the site of ribozyme-mediated cleavage.

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thus the first wash was followed by a short 10-sec rinse. Since longer incubation times (15–60 min) with the WCE did not change the results, the shortest incubation time (15 min) was selected to help reduce the total processing time between the addition of the WCE and the first RNA elution (∼20 min).

The amount of cell extract used for the RNA pull-down assay was found to be very critical. Using 3.5 nmol of SL-let-7g, the addition of ∼300 µg WCE in a final volume of 400 µL was optimal with both NT2 and P19 (mouse embryonal carcinoma; not shown) cells. However, it may be important to adjust the amount of WCE when using either other target RNAs or other cell types. Finally, the spin-cup purification method was found to be one of the most important factors for maximizing the abundance of protein interactors in the SL-let-7g RNA pull-down while minimizing the level of background binding in the mock pull-down. In this respect, the spin-cup purification method was found to be more efficient than the standard batch method, in addition to being faster and more convenient. Although the use of magnetic resin has been previously reported to improve the specificity of RNA pull-downs (Oeffinger 2012), we observed an increase in background binding with the use of magnetic beads (MagneGST, Promega; results not shown) in comparison with the use of nonmagnetic beads retained by spin cups.

Label-free quantitative mass spectrometry

For our riboproteomics studies, RNA pull-downs were performed in biological triplicate with two ARiBo-tagged miRNA stem–loops of the let-7 family, SL-let-7a-1 and SL-let-7g, as well as with two experimental controls, the ARiBo tag alone (mock) and an ARiBo-tagged SL-let-7g variant that retains the duplex region but not the terminal loop (dlet-7g; Fig. 2; Desjardins et al. 2011). The general scheme used for label-free quantitative MS of ARiBo pull-downs is illustrated in Figure 5. Briefly, the sample and control RNA pull-down assays are performed independently with the same NT2 WCE preparation, and the tryptic digests of the eluted samples are analyzed by nano-liquid chromatography coupled with tandem mass-spectrometry (nano-LC–MS/MS). For each ARiBo pull-down, the spectral counts for each protein identified were analyzed using SAINT (Significance Analysis of INTeractome), an algorithm that provides probabilistic scoring of AP-MS data analysis and has been successfully adopted in several studies (Breitkreutz et al. 2010; Skarra et al. 2011; Oeffinger 2012; Tate et al. 2013). The SAINT analysis takes into account data from biological replicates as well as from negative control purifications (Teo et al. 2014). Here, the SAINTexpress scores were computed based on spectral counts of the three individual replicates using either the mock purification (Smock score) or the dlet-7g purification (Sdlet score) as the negative control. SAINT analysis led to the identification of several binding partners for the SL-let-7a-1 and SL-let-7g RNAs using Smock and Sdlet scores ≥0.65 (Table 1; Supplemental Tables S1, S2).

Identification of several common protein interactors for SL-let-7a-1 and SL-let-7g

Quantitative MS analysis allowed the identification of 86 and 50 proteins from the SL-let-7a-1 and SL-let-7g pull-downs, respectively. This includes a set of 42 proteins common to both SL-let-7 RNAs (Fig. 6A; Table 1; Supplemental Table S1). A small subset of these 42 proteins were selected for further validation by Western blot analysis, namely Nucleolin, IGF2BP1 and Lin28B, and confirmed their selective enrichment in the SL-let-7a-1 and SL-let-7g RNA pull-downs in comparison with a mock pull-down (Fig. 6B). Interestingly, these three proteins have been associated with cancer progression (Thornton and Gregory 2012; Bell et al. 2013; Pichierri et al. 2013) and are known to bind RNA sequences that match homologous sequence elements of pre-let-7 RNAs [5′-UUAGG-3′ for Nucleolin (Ishikawa et al. 1993), 5′-(C/A)CA(C/U)-3′ for IGF2BP1 (Patel et al. 2012) and 5′-GGAG-3′ for Lin28B (Graf et al. 2013)], suggesting that they may regulate several members of the let-7 family.

Given that the human mRNA interactome comprises over 1200 RNA-binding proteins likely important for numerous aspects of RNA processing and function (Baltz et al. 2012;
down and the RNA purification have been optimized such that for the RNA pull-down (19 ng of SL-let-7g to be purified (Di Tomasso et al. 2011, 2012a). The protocols for the RNA pull-down are equivalent to a maximum of 50 ng of ARiBo-tagged SL-let-7g from the transcription reaction or standard amounts shown, where 1× corresponds to the volumes of affinity purification eluates (or washes) highlights the purified RNA.

For quantitative analysis, as previously described (Di Tomasso et al. 2011, 2012a). The red box raises the possibility that there may be additional recognition elements for binding than those present in the SL-let-7 RNAs (Ha 2011; Connerty et al. 2015), such as specific elements at the 5’- and 3’-ends (e.g., 5’-monophosphate and 3’-OH). Other proteins involved in miRNA biogenesis, such as Musashi1 (Kawahara et al. 2012), are missing from our analysis most likely due to our experimental conditions, particularly the specific cell type used for preparing the WCE and the type of miRNA stem-loops tested. Surprisingly, Lin28A, which is present in the NT2 cell extract and was identified by Western blot analysis of SL-let-7g pull-downs (Fig. 4B,C), did not pass the cutoff to be considered an interactor of either SL-let-7 RNA in the MS analysis. Based on spectral counts, Lin28A was identified as a highly abundant protein in all RNA pull-downs reported in this study, including the control samples, but SAINTexpress scores of zero were obtained for the analysis of the SL-let-7a-1 and SL-let-7g RNA pull-downs (Supplemental Table S2). Thus, Lin28A is a false negative of the MS analysis likely due to the high abundance of Lin28A in NT2 cells and the ability of this protein to bind nonspecifically to RNA at high concentration. This result raises the possibility that there may be additional false negatives like Lin28A that are not identified in our AP-MS study due to the stringency of the MS analysis criteria.

Interestingly, seven proteins were identified in both the SL-let-7a-1 and SL-let-7g pull-downs when using the mock control but not when using the diet-7g control (Table 1; Fig. 6). These results strongly indicate that these proteins specifically interact with the duplex region of these SL-let-7 RNAs. Although the RNA-binding properties of four of these seven proteins (HMGA1, TCOF1 associated with the Treacher Collins syndrome 1, SON, and HSPE1) have not been well characterized, the other three (STAU1, ILF2, and ILF3) are
known double-stranded RNA-binding proteins (Satoh et al. 1999; Wickham et al. 1999). The ILF2 and ILF3 proteins form the ILF2-ILF3 heteromeric complex, also termed the NF45-NF90 complex, which has been shown to negatively regulate miRNA biogenesis (Sakamoto et al. 2009). To the best of our knowledge, STAU1, more commonly known as Staufen1, has not been directly implicated in miRNA biogenesis (Ren et al. 2016). However, genome-wide analysis of Staufen1-associated mRNAs provides strong evidence that preferential Staufen1 targets for in vivo binding are double-stranded regions that contain mismatches and/or unpaired bases (Kim et al. 2007; Laver et al. 2013), like those found in miRNA stem–loops.

DISCUSSION

Several proteomic studies have previously exploited AP-MS methodologies to identify proteins that interact with miRNAs stem–loops. However, among the 94 proteins identified in our riboproteomic analysis of the SL-let-7a-1 and SL-let-7g pull-downs, only 36 have been previously identified in proteomic studies with the same miRNA stem–loops, including proteomic studies using the same NT2 cell line (Table 1). Therefore, our AP-MS method allows for the identification of several SL-let-7-associating proteins that were not previously identified and, given the stringency of the analysis, are likely to represent genuine interactors of let-7 miRNA stem–loops. Some of these newly identified SL-let-7 interactors are known to be involved in important biological processes, such as development and tumorigenesis. Interestingly, we find all three IGF2BP human paralogs, IGF2BP1, IGF2BP2, and IGF2BP3, which have been implicated in development, metabolic control and tumorigenesis and are known to direct mRNA stabilization, translation, and transport (Bell et al. 2013).

Similarly, they could regulate miRNA biogenesis and transport of immature forms of miRNAs. We also identified five proteins, IGF2BP1, HNRNPU, SYNCRIP, YBX1, and DHX9, that constitute the coding region determinant (CRD)-mediated complex known to promote mRNA stability of the c-myc oncogene (Weidensdorfer et al. 2009), suggesting that the CRD-mediated complex could also play a role in stabilizing immature forms of miRNAs. Future studies are needed to validate that the proteins identified here directly bind to the SL-let-7a-1 and SL-let-7g RNAs in the absence of the ARiBo tag and to establish whether or not these proteins regulate miRNA biogenesis. In addition, it will be interesting to determine if these proteins interact with other miRNA stem–loops since only a handful of pre-miRNAs have been characterized by AP-MS to date.

As in other AP-MS studies, the use of a tag for affinity purification may prevent the binding of proteins that would
TABLE 1. Partial MS analysis results listing proteins identified in both the SL-let-7a-1 and the SL-let-7g ARiBo pull-downs

| Gene name | Protein name | Uniprot entry | SL-let-7a-1 | SL-let-7g |
|-----------|--------------|---------------|------------|----------|
|           |              |               | Versus Mock | Versus dlet-7g | Versus Mock | Versus dlet-7g | RBP | 7a-1 | 7g |
| C7orf50   | Uncharacterized protein C7orf50 | Q9BRJ6 | ○ | ○ | ○ | ○ | • |        |
| CSDAP1_YBX3 | Y-box-binding protein 3 | P16989 | ○ | ○ | ○ | ○ | • | 4 |
| CSNK2A1   | Casein kinase II subunit α | P68400 | ○ | ○ | ○ | ○ | • |        |
| DDX5      | Probable ATP-dependent RNA helicase DDX5 | P17844 | ○ | ○ | ○ | ○ | • | 3, 4 | 6 |
| DHX9      | ATP-dependent RNA helicase A | Q08211 | ○ | ○ | ○ | ○ | • | 1,3,4,5 | 6 |
| CRSF1     | G-rich sequence factor 1 | Q12849 | ○ | ○ | ○ | • |        |
| HMGA1     | High-mobility group protein HMGA1 | P17096 | ○ | ○ | ○ | • |        |
| HNRNPA0   | Heterogeneous nuclear ribonucleoprotein A0 | Q13151 | ○ | ○ | ○ | • | 1,3,4,5 | 6 |
| HNRNPA3   | Heterogeneous nuclear ribonucleoprotein A3 | P51991 | ○ | ○ | ○ | ○ | • | 3, 4 | 6 |
| HNRNPA8   | Heterogeneous nuclear ribonucleoprotein A/B | Q99729 | ○ | ○ | ○ | • | 4 | 6 |
| HNRNPD    | Heterogeneous nuclear ribonucleoprotein D0 | Q14103 | ○ | ○ | ○ | • | 3 |        |
| HNRNPU    | Heterogeneous nuclear ribonucleoprotein U-like protein 1 | Q980839 | ○ | ○ | ○ | • | 3, 4 | 6 |
| HSPE1     | 101 kDa heat shock protein, mitochondrial | P61604 | ○ | ○ | • |        |
| IGF2BP1   | Insulin-like growth factor 2 mRNA-binding protein 1 | Q9NZI8 | ○ | ○ | ○ | • | 3, 4 | 6 |
| IGF2BP2   | Insulin-like growth factor 2 mRNA-binding protein 2 | Q9Y6M1 | ○ | ○ | ○ | • |        |
| IGF2BP3   | Insulin-like growth factor 2 mRNA-binding protein 3 | O00425 | ○ | ○ | ○ | • |        |
| ILF2      | Interleukin enhancer-binding factor 2 | Q12905 | ○ | ○ | • | 3, 4, 5 | 6 |
| ILF3      | Interleukin enhancer-binding factor 3 | Q12906 | ○ | ○ | • | 1,3,4,5 | 6 |
| LIN28B    | Protein Lin-28 homolog B | Q6ZN17 | ○ | ○ | ○ | • | 1, 3, 4, 5 | 6 |
| NCL       | Nucleolin | P19338 | ○ | ○ | ○ | • | 2,4,5 |        |
| NONO      | Non-POU domain-containing octamer-binding protein | Q15233 | ○ | ○ | • | 4 | 6 |
| NSUN2     | tRNA [cytosine(34)-C(5)]-methyltransferase | Q08J23 | ○ | ○ | ○ | • |        |
| RPL22     | 60S ribosomal protein L22 | P35268 | ○ | ○ | ○ | • |        |
| RPS17;RPS17L | 40S ribosomal protein S17 | P08708 | ○ | ○ | ○ | • |        |
| RPS7      | 40S ribosomal protein S7 | P62081 | ○ | ○ | ○ | • |        |
| SON       | Protein SON | P18583 | ○ | ○ | • |        |
| SRP14     | Signal recognition particle 14 kDa protein | P37108 | ○ | ○ | • |        |
| SRSF1     | Serine/arginine-rich splicing factor 1 | Q07955 | ○ | ○ | • | 3 | 6 |
| SRSF9     | Serine/arginine-rich splicing factor 9 | Q13242 | ○ | ○ | • | 3 |        |
| SSB       | Lupus La protein | P05455 | ○ | ○ | • | 3 |        |
| STAU1     | Double-stranded RNA-binding protein Staufen homolog 1 | O95793 | ○ | ○ | • | 3 | 4, 6 |
| STRBP     | Spermatid perinuclear RNA-binding protein | Q9659 | ○ | ○ | • | 3, 4 | 6 |
| SUB1      | Activated RNA polymerase II transcriptional coactivator p15 | P53999 | ○ | ○ | • |        |
| SYNCRIP   | Heterogeneous nuclear ribonucleoprotein Q | Q96050 | ○ | ○ | • |        |
| TCOF1     | Treacle protein | Q13428 | ○ | ○ | • |        |
| TFAM      | Transcription factor A, mitochondrial | Q00059 | ○ | ○ | • |        |
| THOC4;ALYREF | THO complex subunit 4 | Q866V1 | ○ | ○ | • | 6 |        |
| TRMT61A   | tRNA [adenine(58)-N(1)]-methyltransferase catalytic subunit | Q96FX7 | ○ | ○ | • |        |
| XRCC6     | X-ray repair cross-complementing protein 6 | Q12956 | ○ | ○ | • |        |
| YBX1      | Nuclease-sensitive element-binding protein 1 | P67809 | ○ | ○ | • |        |
| ZC3HA1    | Zinc finger CCCH-type antiviral protein 1 | Q722W4 | ○ | ○ | • |        |

Positive identification of a protein interactor (open circle) from the SL-let-7a-1 and SL-let-7g pull-downs is based on a SAINTexpress score ≥0.65. Results are listed using either the mock or the dlet-7g ARiBo pull-down as the negative control. RNA-binding proteins (RBPs; black dot) are inferred based on recent genome-wide AP-MS studies of mRNA-binding proteins (Baltz et al. 2012; Castello et al. 2012; Kwon et al. 2013; Beckmann et al. 2015). Results from previous AP-MS studies are provided for let-7a-1 stem–loops (7a-1; 1: WCE from Huh7 cells [Heo et al. 2008]; 2: with nuclear extract from HeLa cells [Michlewski et al. 2008; Michlewski and Caceres 2010]; 3: WCE from Lin28A-transfected HEK293T cells [Heo et al. 2009]; 4: WCE from P9 cells [Chang et al. 2013]; and 5: WCE from HeLa and NT2 cells [Lee et al. 2013]) and a let-7g stem–loop (7g; 6: with P19 cells [Viswanathan et al. 2008]).

The SAINTexpress scores for the identification of the YBX1 protein from the SL-let-7a-1 and SL-let-7g pull-down versus the mock control are just below the cutoff (0.65 > score ≥ 0.60).
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FIGURE 6. Summary and validation of the AP-MS results for SL-let-7a-1 and SL-let-7g. (A) The overlap of identified proteins in the SL-let-7a-1 and SL-let-7g ARiBo pull-downs is shown using Venn diagrams. The total proteins identified were taken from proteomic analysis using the mock and dlet-7g controls. The proteins that likely bind the duplex region are those identified using the mock control but not the dlet-7g control. (B) Validation of the AP-MS results by Western blots of the elution fractions (50% of E1) from the SL-let-7a-1, SL-let-7g, and mock pull-downs.

normally interact with the free target or, inversely, induce the binding of proteins that do not normally bind to the free target. In particular, a common concern when using an RNA tag is the possibility for interactions between the tag and the target RNA that would disrupt the folding of the target RNA. Using the activatable ARiBo tag, one can quickly verify that this tag does not induce significant misfolding of the target RNA. In the present study, this was done by testing the efficiency of cleavage of the ARiBo tag from the fusion RNAs in the presence of 1 mM GlcN6P (lanes 15–18 in Fig. 3A,B). For both the SL-let-7a-1-ARiBo (not shown) and SL-let-7g-ARiBo (Fig. 3) fusion RNAs, efficient cleavage of the ARiBo tag was observed under these conditions, providing clear evidence that the ARiBo tag and target RNA fold independently of each other within the fusion RNAs.

Our AP-MS approach is particularly powerful given that the ARiBo pull-down significantly minimizes nonspecific binding (low protein levels in the mock control; Fig. 4A), and the MS analysis provides a stringent quantitative assessment of true protein interactors. The high sensitivity and specificity of the RNA pull-down results in part from the use of optimized conditions for the RNA pull-downs, including the amount of cell extract, buffer conditions, and elution conditions, but also depends on the use of spin-cup devices during the purification to efficiently wash protein contaminants without losing resin. The ARiBo tag itself contributes significantly to the high sensitivity and specificity of the RNA pull-down by affording high-affinity immobilization via the λboxB/N peptide complex (Koff of 10 pM) (Austin et al. 2002; Di Tomasso et al. 2011) as well as highly efficient elution of the RNA–protein complex by activatable cleavage of the glmS ribozyme. Similarly, activatable cleavage of an RNA tag has previously been exploited for AP-MS studies using the activatable Cas4 enzyme of a cognate RNA substrate tag, where it was shown to dramatically reduce the amount of background protein contaminants (Lee et al. 2013). In comparison to a noncleavable system, elution by cleavage of the target RNA from the immobilized tag prevents the elution of proteins that are bound to the tag and thus likely reduces the amount of eluted proteins that are not specifically associated with the target RNA. Moreover, since cleavage of the ARiBo tag is activated by a small metabolite (Glc6NP), this prevents contamination of the eluate by a larger molecule that could potentially interfere with the MS analysis. An important advantage of the activatable glmS ribozyme compared to other cleavable tags is that efficient cleavage keeps the RNA elution period to a minimum (15 min) such that the RNA pull-down can be performed very quickly under conditions that help maintain the integrity of the RNA–protein complexes. Thus, the ARiBo pull-down procedure is also well suited for other applications, such as the purification of RNA–protein assemblies for biochemical and structural characterization. Since it can be performed quickly and efficiently, the ARiBo pull-down could also be used to streamline high-throughput isolation of proteins that associate with target RNAs.

In summary, combining the ARiBo pull-down method with label-free quantitative mass spectrometry has allowed us to better define the protein interactome of the stem–loops of two specific miRNAs, human let-7a-1 and let-7g. Our method is well suited to investigate the protein interactors of other interesting RNAs found in living cells so that we can better understand how these RNAs collaborate with proteins to sustain a wide range of biological processes.

MATERIALS AND METHODS

Cell culture and whole cell extract (WCE) preparation

The human pluripotent embryonal carcinoma cell line NTERA-2 cl. D1 (NT2; ATCC) was maintained in Dulbecco’s modified Eagle’s Medium (DMEM; Wisent) supplemented with 10% (v/v) fetal bovine serum (FBS; Wisent) at 37°C and in 10% CO2. NT2 cells grown to near confluency in 150-mm Petri dishes were washed once with cold PBS (10 mM Na2HPO4, 2 mM KH2PO4, 2.7 mM KCl, and 140 mM NaCl, adjusted to pH 7.4) and harvested by scraping. The resuspended cells were washed twice with cold PBS, and the cell debris removed by centrifugation. The pellet was incubated for 20 min on ice with WCE lysis buffer (50 mM HEPES [pH 7.5], 100 mM NaCl, 1% NP-40 and 1 mM DTT) supplemented with protease inhibitors (0.1 mM AEBSF, 1 μg/mL aprotinin, 1 μM pepstatin, and 1 μg/mL leupeptin), vortexing every 5 min. The cell lysate was then clarified by centrifugation for 10 min at 16,000g. The resulting WCE was snap frozen in a dry-ice/ethanol bath and kept at −80°C until use. The protein concentration of the WCE was quantified with a Micro BCA protein assay kit (Pierce).

Recombinant protein expression and purification

Previously established procedures were employed for the expression and purification of the LIN-L+-GST fusion protein (Di Tomasso et al. 2012). The resulting LIN28B N+-L+-GST fusion protein (Di Tomasso et al. 2012) was modified Eagle’s Medium (DMEM; Wisent) supplemented with 10% (v/v) fetal bovine serum (FBS; Wisent) at 37°C and in 10% CO2. NT2 cells grown to near confluency in 150-mm Petri dishes were washed once with cold PBS (10 mM Na2HPO4, 2 mM KH2PO4, 2.7 mM KCl, and 140 mM NaCl, adjusted to pH 7.4) and harvested by scraping. The resuspended cells were washed twice with cold PBS, and the cell debris removed by centrifugation. The pellet was incubated for 20 min on ice with WCE lysis buffer (50 mM HEPES [pH 7.5], 100 mM NaCl, 1% NP-40 and 1 mM DTT) supplemented with protease inhibitors (0.1 mM AEBSF, 1 μg/mL aprotinin, 1 μM pepstatin, and 1 μg/mL leupeptin), vortexing every 5 min. The cell lysate was then clarified by centrifugation for 10 min at 16,000g. The resulting WCE was snap frozen in a dry-ice/ethanol bath and kept at −80°C until use. The protein concentration of the WCE was quantified with a Micro BCA protein assay kit (Pierce).
et al. 2011, 2012b) and the human Lin2A protein (Desjardins et al. 2011).

In vitro transcription of ARiBo-tagged RNAs and affinity purifications

All plasmids for in vitro transcription of ARiBo-tagged RNAs were obtained by inserting double-stranded DNA fragments generated by recursive PCR into the HindIII/Apal sites of the pARiBo+ plasmid (Bowman et al. 2012; Salvail-Lacoste et al. 2013), and the integrity of each plasmid was verified by sequence analysis. The ARiBo-tagged RNAs were transcribed for 3 h at 37°C using the following reaction conditions: 40 mM HEPES (pH 8.0), 50 mM DTT, 0.1% Triton X-100, 1 mM spermidine, 4 mM of each NTP (ATP, CTP, UTP, and GTP), 25 mM MgCl2, 60 µg/mL His-tagged T7 RNA polymerase, 3 U/mL RNAse in ribonuclease inhibitor (Promega), and 80 µg/mL linearized plasmid DNA template (Di Tomasso et al. 2012a). Transcription reactions were stopped by adding 25 mM EDTA (pH 8.0) and then stored at −20°C. Cleavage of the ARiBo tag in the transcription reaction (glmS cleavage) and batch-affinity purifications were performed as previously described (Di Tomasso et al. 2014). In vitro transcriptions, glmS cleavage reactions and batch-affinity purifications were monitored by running aliquots on 7.5% denaturing gels, staining with SYBR-Gold (Invitrogen) and affinity purifications were monitored by running aliquots on 7.5% denaturing gels, staining with SYBR-Gold (Invitrogen) and subsequent fluorescence detection on a Bio-Rad ChemiDoc MP Imaging system (Di Tomasso et al. 2014).

ARiBo pull-down assay

RNA-associated proteins were isolated with a modified version of the ARiBo affinity purification protocol (Di Tomasso et al. 2011, 2012a, 2014). Briefly, 17.5 nmol of LN−L−-GST protein was first added to an aliquot of the transcription reaction (30–200 µL) containing 3.5 nmol of ARiBo-tagged RNA, and the volume was completed to 400 µL with WCE equilibration buffer (50 mM HEPES, pH 7.5, 100 mM NaCl, and 5 mM MgCl₂). After a 15-min incubation with rotation at room temperature (RT), the RNA–protein mix was added to a spin-cup column (Pierce) containing 125 µL of washed GSH-Sepharose 4B resin (from 163 µL of slurry; GE Healthcare) and incubated for another 15 min at RT. The load eluate was collected by centrifugation for 1 min at 5000g. The resin was washed once with 400 µL of WCE equilibration buffer. Unless specified otherwise, all washes included incubation for 5 min at RT with rotation and centrifugation for 1 min at 5000g. After this first wash (W1), 300 µg of NT2 WCE diluted in WCE equilibration buffer (400 µL total volume) was loaded to the RNA-coupled resin and incubated for 15 min at 4°C with rotation, followed by centrifugation. The resin was then washed once (W2) and rinsed once (six tube inversions followed by centrifugation) with 400 µL of WCE equilibration buffer. For the first elution of RNA and associated proteins (E1), the resin was incubated in 400 µL of elution buffer (20 mM Tris [pH 7.6], 10 mM MgCl₂, and 1 mM GlnN6P) for 15 min at 37°C and then for 5 min at RT prior to centrifugation. For the second elution (E2), the resin was washed once with 400 µL of WCE equilibration buffer. Finally, the resin was cleared with 400 µL of 2.5 M NaCl. Generally, 10% aliquots of the load, washes and elution samples (LE, W1, WCE, W2, E1, E2, NaCl) were kept for analysis by denaturing gel electrophoresis (Di Tomasso et al. 2011, 2012a) and Western blot. The rest of the first elution sample was lyophilized for analysis by either Coomassie-stained SDS-PAGE or mass spectrometry. RNA pull-downs were performed in triplicate for all RNAs (the ARiBo-tagged SL-let-7a-1, SL-let-7g, and dlet-7 as well as the mock control) using three different WCE preparations prepared from separate batches of NT2 cells.

Coomassie-stained SDS-PAGE and Western blotting

A fraction of the first elution sample was precipitated with 25% (v/v) trichloroacetic acid. The sample was vortexed, incubated for 30 min on ice, and centrifuged for 20 min at 16,000g. The pellet was then washed two times by adding 200 µL of acetone and centrifuging 15 min at 16,000g. Following the drying of the pellet for 5 min in a 95°C dry bath, the samples were resuspended in Laemml SDS-PAGE sample buffer and run on a 12% SDS-PAGE. The gel was either stained with Coomassie Blue or transferred to an Immun-Blot PVDF membrane (Bio-Rad) for Western blot analysis using Lin28A (Millipore #07-1385), Lin28B (Cell Signaling Technology #4186), IFG2BP1 (Abcam #ab107205), or nucleolin (Abcam #ab134164) antibodies. A ChemiDoc MP Imaging system was used for chemiluminescence detection, and band intensities were quantified using the ImageLab software version 4.0.1 (Bio-Rad).

Mass spectrometry data acquisition and analysis

Proteins eluted following RNA pull-down assays were solubilized in 6 M urea and 100 mM ammonium bicarbonate. Cysteine residues were reduced with 10 mM DTT and alkylated with 55 mM iodoacetamide. The samples were then diluted to reduce the urea concentration to <1 M and the proteins in-solution digested with trypsin for 16 h at 37°C. The reactions were quenched by acidification with trifluoroacetic acid and passed over a C18 spin column (Pierce) to remove nonpolar contaminants, including the RNA and salts. The purified peptides were solubilized in 2% acetonitrile and 0.5% acetic acid for online mass spectrometry analysis using an Orbitrap Fusion (Thermo) mass spectrometer. Peptides were trapped on a trapping column and separated on a 75 µm × 15 cm, 2 μm Acclaim PepMap reverse phase column (Thermo Scientific) using an UltiMate 3000 RSLCnano HPLC (Thermo Scientific) at a flow rate of 300 nL/min. Peptides were eluted into the mass spectrometer using a 120-min linear gradient from 96% mobile phase A (0.1% formic acid) to 55% mobile phase B (80% acetonitrile and 0.08% formic acid). Parent full-scan mass spectra were collected in the Orbitrap mass analyzer set to acquire data at 120,000 FWHM resolution; ions were then isolated in the quadrupole mass filter, fragmented within the HCD cell (HCD normalized energy 32%, stepped ±3%), and the product ions were analyzed in the ion trap. Monoisotopic precursor selection and dynamic exclusion, with a 30-sec duration and a 10-ppm mass tolerance, were enabled. Proteome Discoverer 1.4 (Thermo) was used to search the data against human proteins from the NCBI nr database using SequestHT and MSAmanda (Dorfer et al. 2014). The sequence of the LN−L−-GST fusion protein was manually added to the database to allow its identification. The search was limited to tryptic peptides, with maximally two missed cleavages allowed. Cysteine carboxamidomethylation was set as a fixed modification; methionine oxidation was allowed as a variable modification. The precursor mass tolerance was 10 ppm, and the fragment mass tolerance was 0.6 Da. The Percolator node was used to score and rank peptide matches at a 1% false discovery rate.

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rate using a reversed algorithm. Proteins were required to be represented by at least two unique peptide sequences. Statistical comparison of the interactome from biological triplicate experiments was performed using the SAINtExpress algorithm (Teo et al. 2014) based on spectral counts and using default settings. To be considered as interactors, proteins had to be enriched at least twofold as compared with the control samples and have a SAINtExpress score $\geq 0.65$. The selected cutoff score of 0.65 ensures that the interaction has been detected with high probability in at least two of the three replicates.

**SUPPLEMENTAL MATERIAL**

Supplemental material is available for this article.

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