Comparative analysis of LIN28-RNA binding sites identified at single nucleotide resolution

Elizabeth Ransey, Anders Björkboom, Victor S. Lelyveld, Przemysław Biecek, Lorena Pantano, Jack W. Szostak, and Piotr S Liz

Introduction

LIN28 is a highly conserved RNA-binding protein that was first described as the product of the heterochronic gene, LIN28, in C. elegans. In mammals, the two LIN28 paralogs, LIN28A and LIN28B, play roles in a wide range of cellular processes, including stem cell self-renewal and pluripotency, skeletal myogenesis, glucose metabolism in diabetes, and tissue repair. These proteins are upregulated in ~15% of human tumors and cancer cell lines, and elevated expression is associated with a poor prognosis and increased aggression in numerous malignancies, including germ cell tumors, colon cancer, and ovarian cancer. Signaling components in the Wnt pathway have been shown to cooperate with LIN28 to increase the severity and invasiveness of colorectal cancer. LIN28B also promotes metastasis of colon cancer and tumorigenesis of the intestinal epithelium.

LIN28 exerts its profound phenotypic effects by acting as a negative regulator of let-7 miRNA biogenesis. Specifically, LIN28 binds let-7 precursors and prevents miRNA maturation, limiting cellular differentiation. High-resolution crystal structures of mouse LIN28A-let-7 complexes and other structural and biochemical data revealed that LIN28 inhibits Dicer processing through steric hindrance and by locally unwinding the cleavage site. Furthermore, LIN28 proteins recruit the terminal uridylyltransferase, TUT4, to uridylate bound miRNAs, resulting in degradation by the Dis3l2 exonuclease.

Recent developments suggest that LIN28 also functions through let-7 independent mechanisms. Several studies using crosslinking and immunoprecipitation with deep sequencing (CLIP-seq) and photoactivatable ribonucleoside analog CLIP (PAR-CLIP) have identified thousands of potential pre-mRNA or mRNA targets of LIN28. These methods rely on irradiation by UV light to generate covalent RNA-protein heterocjugates in live cells, allowing for the isolation of RNA binding proteins (RBPs) by immunoprecipitation and subsequent high-throughput sequencing of crosslinked RNAs. CLIP was originally developed to address limitations of non-covalent RBP bioinformatics and sequencing analyses suggest guanine in mammalian cells, MS/MS identifies uridine. This work suggests the need for comprehensive analysis and validation of crosslinking methodologies.

ABSTRACT

It remains a formidable challenge to characterize the diverse complexes of RNA binding proteins and their targets. While crosslink and immunoprecipitation (CLIP) methods are powerful techniques that identify RNA targets on a global scale, the resolution and consistency of these methods is a matter of debate. Here we present a comparative analysis of LIN28-pre-let-7 UV-induced crosslinking using a tandem mass spectrometry (MS/MS) and deep sequencing interrogation of in vitro crosslinked complexes. Interestingly, analyses by the two methods diverge in their identification of crosslinked nucleotide identity – whereas bioinformatics and sequencing analyses suggest guanine in mammalian cells, MS/MS identifies uridine. This work suggests the need for comprehensive analysis and validation of crosslinking methodologies.

CONTACT Piotr S Liz sliz@hms.harvard.edu 240 Longwood Avenue, Department of Biological Chemistry and Molecular Pharmacology and Pediatrics, Harvard Medical School, Boston, MA, 02115, USA. Supplemental data for this article can be accessed on the publisher’s website. © 2017 Elizabeth Ransey, Anders Björkboom, Victor S. Lelyveld, Przemysław Biecek, Lorena Pantano, Jack W. Szostak and Piotr S Liz. Published with license by Taylor & Francis Group, LLC.

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containing the sequence GGAG, the signature LIN28 recognition element present in let-7 miRNAs. Furthermore, one report identified an enrichment of pyrimidine-rich binding motifs, consistent with sequences recognized by the LIN28 cold shock domain (CSD) in let-7 targets, suggesting that LIN28 interactions with mRNA may have binding determinants that mirror those previously identified in let-7 miRNA targets.

Building on CLIP methods, crosslink induced mutation site (CIMS) analysis has emerged as a powerful bioinformatic tool for the elucidation of single nucleotide resolution crosslink interaction information derived from CLIP-seq data sets. This method aims to identify mutations, primarily deletions and substitutions, that occur during reverse transcription at presumed crosslink sites within CLIP reads. Despite the availability of numerous LIN28 CLIP-seq and PAR-CLIP studies, crosslink profiles from mutational analysis have been reported in only one study (see ref. 24), which found an increased prevalence of mutations at guanine residues that were apparently localized within a LIN28 GGAG recognition motif. To our knowledge, no other CLIPS studies have been examined by CIMS, nor has the crosslinked side chain on LIN28 been concurrently elucidated.

In this work, we endeavored to characterize the products of in vitro UV-induced crosslinking within a well-characterized complex of LIN28A and a precursor let-7 fragment. Specifically, we used mass spectrometry (MS) to identify the discrete position of a crosslinked interaction on both the protein and the RNA components simultaneously. Furthermore, we performed sequencing analyses of the crosslinked RNA component to identify any crosslink-induced mutations (CIMS). Interestingly, we determined that the two methods non-redundantly determined different crosslink sites and crosslinked nucleotide identities. Overall, our work suggests that high-precision analysis methods of RNA-protein crosslinks must be cross-validated to avoid methodology-specific conclusions and gain comprehensive information concerning crosslinked interactions.

Results

Recombinant model LIN28A and pre-let-7f complexes can be crosslinked in vitro

To evaluate the in vitro protein-RNA crosslinking profile of a representative bipartite LIN28A-RNA complex, we generated complexes comprised of a previously reported, truncated mouse LIN28 construct (LIN28A-ΔΔ) and a correspondingly modified pre-element let-7f miRNA substrate, (preFm,let-7f) (Fig. 1A, B). LIN28A-ΔΔ consists of amino acids D33-K187 of the full-length protein and lacks the random coil N- and C-termini as well as a nine amino acid internal flexible linker between the CSD and zinc-knuckle domain (ZKD) (Fig. 1A). PreFm,let-7f has a 5-nucleotide deletion between the AYYYH (the CSD-binding pyrimidine-rich sequence motif, where Y = C or U and H = A, C, or U) and GGAG elements to accommodate the decreased space between the LIN28A binding domains (Fig. 1B). These truncated components were previously crystallized as a complex, which reflected the interactions between the wild type full-length LIN28A and preE-let-7f, as determined by functional studies. The binding affinity between LIN28A-ΔΔ and preFm,let-7f was comparable to the full-length LIN28A affinity for its corresponding preE-let-7f (47 – 190 nM) (Fig. 1C), in agreement with previous data. We also observed that purified LIN28A-ΔΔ:preFm,let-7f complexes could be crosslinked with comparable efficiency as full-length LIN28A:preE-let-7f (Fig. 1D). This finding suggests that in vitro complexes that incorporate these truncated components are sufficient to mimic in vivo binding of the native protein with this miRNA intermediate.

With the aim of determining crosslink sites via MS and CIMS analyses, we established a workflow to generate crosslinked heteroconjugate samples composed of the aforementioned truncated constructs. Complexes of LIN28A-ΔΔ and preFm,let-7f were exposed to 254 nm UV light, and crosslinked complexes were isolated and trypsinized to yield samples of peptide-modified RNA. The peptide-modified RNA is suitable for both liquid chromatography-mass spectrometry (LC-MS) and sequencing analyses. Subsequent RNA digestion by formic acid (FA) rendered the sample suitable for tandem mass spectrometry (MS/MS) (Fig. 1E).

Mass spectrometry reveals a crosslinking site between the let-7 pre-element terminal loop and LIN28A cold-shock domain

To assign discrete crosslink sites in model recombinant RNA-LIN28 complexes, we used an RNA-directed enrichment approach and mass spectrometry. The full-length, unmodified preFm,let-7f RNA has an exact calculated mass of 8052.0600 Da, and crosslinked heteroconjugates were initially sought by scrutinizing species with neutral mass gains that might correspond to tryptic peptide addition. Initially, we observed a discrete species with a low resolution mass of 9268 Da, consistent with a mass gain corresponding to the predicted tryptic fragment MGFGFSLMTAR (residues 51 – 61 in full length LIN28).

To examine this specific peptide-RNA heteroconjugate, enriched peptide-RNA samples were stringently hydrolyzed in 50% (v/v) formic acid (FA) at 80°C for 2 h. Survey spectra were searched for peptide-nucleotide heteroconjugates as either simple mass neutral conjugates or those exhibiting neutral mass losses that might correspond to tryptic peptide addition. Initially, we observed a discrete species with a low resolution mass of 9268 Da, consistent with a mass gain corresponding to the predicted tryptic fragment MGFGFSLMTAR (residues 51 – 61 in full length LIN28).

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y7 – y10 but not for smaller products, indicating that the UMP is crosslinked to Phe55 in LIN28A-ΔΔ.

To assign the crosslinked position within the pre-element RNA, we sought to observe crosslinked heteroconjugates composed of peptides covalently bound to larger nucleotide species: either di- or tri-nucleotides. Crosslinked heteroconjugates were generated and hydrolyzed as before, with the exception that the acid digest time was reduced (30 min) and temperature lowered.
Table 1. Masses and nucleotide composition overlap of crosslinked mono-, di- and tri-nucleotides that identify U11 as the most probable crosslinking counterpart to Phe55.

| Species      | Mass (Da) | RNA Sequence          |
|--------------|-----------|-----------------------|
| M + preEM-let-7f | 9268      | GGGGUAGUGAU, UUUACCUCUGAGAU |
| M + GAU       | 2214.7149 | GAU                   |
| M + AUU       | 2175.6949 | AUU                   |
| M + UUU       | 2152.6627 | UUU                   |
| M + AU        | 1869.6682 | AU                    |
| M + U         | 1846.6384 | U                     |
| M + UMP       | 1540.6120 |                       |

(60°C) to prevent complete RNA hydrolysis. From these samples we were able to identify ions corresponding to the same peptide, MGFGFLSMTAR, neutrally conjugated to UMP, as well as to RNA fragments with the following nucleotide compositions: UU, AU, UUU, AUU and GAU (Table 1). We further confirmed the nucleotide composition and peptide sequence of these species using tandem MS (Supp. Fig. 1). Analyzing the composition of these species revealed only one consistent overlapping site in the pre-element RNA sequence: uridine-11 (Table 1). We concurrently confirmed the identity of U11 using a newly developed RNA site-specific stable isotope labeling technique.38 Using pre-element RNA labeled with synthetic isotopes at either the U11 or U12 positions, we observed a mass-shifted isotope distribution exclusively for heteroconjugates arising from complexes formed from RNA labeled at U11. The identified tryptic peptide corresponds to a region within the LIN28A CSD at its binding interface with preEM-let-7f (Fig. 3A), as observed in a high-resolution crystal structure (PDB ID: 3TS0).18 Within this interface, Phe55 is oriented such that the side chain is (at closest proximity) within 3.5 Å of the uracil moiety of U11 within the pre-element terminal loop, giving a planar angle of ~7.5 degrees between the two aromatic rings (Fig. 3A). This contact is consistent with a strong π-π interaction between the two residues and suggests that the mass neutral crosslink identified here is physiologic.

To validate the identified crosslinking site and characterize its contribution to substrate binding, we measured in vitro binding affinities of single point mutants of the determined crosslinking site, Phe55, within the LIN28A-ΔΔ construct. Using gel shift binding assays, we found that a single conservative mutation of Phe55 to tyrosine (F55Y) had a minimal effect on binding (K_D: 100 – 200 nM), whereas an alanine mutation at the same position (F55A) resulted in a significant decrease in affinity (K_D: ~700 nM) (Fig. 3B). These results demonstrate significant contribution of Phe55 to LIN28-RNA binding affinity, even though its contribution to binding specificity is small.18 Despite the range of observed affinities, SDS-PAGE experiments confirmed that both mutant constructs were able to crosslink preEM-let-7f, though to varying extents (Fig. 3C), suggesting the presence of crosslink sites undetected by MS.

**CIMS analysis identifies guanine mutations in CLIP-seq data sets and within in vitro crosslinked LIN28-let-7 complexes**

RNA-protein UV crosslinking causes observable mutations in CLIP sequencing reads, which are presumed to be indicative of crosslinking sites and can be mapped using CIMS analysis.36,37 Thus, we generated a data processing workflow modified from previous CIMS protocols (see ref. 36 and 37) and validated our method by reanalyzing two mouse LIN28A CLIP data sets for which CIMS mutational profiles were reported (see ref. 24). Consistent with that work, our analysis of the monoclonal 35L33G and polyclonal antibody CLIP data sets showed that mutations arose most frequently at guanines (Fig. 4A). Though we observed similar mutation identities and positions, our frequencies were lower, likely due to differences in filtering parameters. Nonetheless, we determined guanines make up 64% and 76% of substitution sites and 43% and 47% of deletion sites for monoclonal and polyclonal antibody data sets, respectively (Fig. 4A).

![Figure 3](image-url) Crosslinking of mutant LIN28 suggests the presence of crosslink sites undetected by MS. (A) Front view surface and side view cartoon representations of the known structure of LIN28A-ΔΔ CSD (green) complexed with preEM-let-7f (gray) (PDB ID: 3TS0). Phe55, the identified crosslink site, contacts U11 of the preEM-let-7f terminal loop. The tryptic peptide MGFGFLSMTAR and Phe55 side chain are highlighted in maroon and U11 nucleotide is highlighted in red. (B) Gel shift binding assays with radiolabeled preEM-let-7f probe, mixed with increasing concentrations of LIN28A-ΔΔ mutant constructs: F55Y (0, 100, 200, 400, 800 nM, 1.6, 3.2 μM) and F55A (0, 22, 44, 180, 700 nM, 2.8 μM). (C) Corresponding SDS-PAGE gels show crosslinked complex bands following UV irradiation.
To examine the consistency of this observation across published LIN28 crosslinking studies, we applied our validated analysis to four published data sets for which crosslink-induced mutations have not yet been reported.23,27 These include two data sets denoted as LIN28A and LIN28-V5 from, ref. 23 (H. sapiens). WormSmall and WormBig (C. elegans) are from ref. 27. Selected data sets include only CLIP experiments performed in tissue culture in the absence of photoactivatable ribonucleoside analogs (PAR-CLIP), to avoid well-known mutation bias. Percentage numbers and colors within frequency plots indicate dominant nucleotide identity and relative enrichment.

We observed variability in crosslink-induced mutation enrichment between these 4 data sets (Fig. 4A). While the two human LIN28A CLIP data sets showed enrichments for substitutions at guanine residues, no nucleotide dominated deletion mutations. Additionally, our analysis of the C. elegans data set showed that uridine nucleotides were the most frequent points of mutation. Though the enrichment of substitutions occurring at uridines was slight, making up 31% and 32% of mutations for the two data sets, we noted a significant enrichment of
uridines as the sites of deletions, which represent 56% and 57% of that type of mutation. Importantly, despite species, group and perhaps quality differences, all data sets were processed using the same analytical workflow which is likely the reason that some frequency plots appear to be irregular.

Since CIMS analysis of endogenous complexes and MS/MS of recombinant complexes point to distinct non-overlapping crosslink sites, we deep sequenced and performed CIMS analysis on recombinant complexes crosslinked in vitro. To determine relative frequencies of mutations arising from protein-RNA crosslinking, we deep sequenced and analyzed three distinct preparations of preEM-let-7f: preEM-let-7f crosslinked with recombinant LIN28A-ΔΔ, UV-exposed preEM-let-7f in the absence of protein, as well as unexposed, unmodified preEM-let-7f RNA (Fig. 4).

Our analysis revealed numerous points of mutation across the RNA sequence that are statistically significant (p < 0.05) (Supp. Table 1). Several sites, however, were mutated in all three RNA samples, including RNA that had no UV exposure (i.e., A22 and A24). Furthermore, a subset of sites were mutated in both the peptide-modified RNA (Fig. 4B, top panel) and the UV-exposed RNA (Fig. 4B, middle panel), indicating that these sites are artifacts of UV exposure and not specifically associated with protein crosslinking (i.e., C16, C17 and C18). Finally, mutation frequencies at positions G1, G2 and G3 were excluded from analytical considerations as, for reasons unclear, several orders of magnitude fewer reads could be mapped to those positions in all samples (Fig. 4B and Supp. Table 1). One major cluster of mutations that was unique to peptide-modified preEM-let-7f was found at positions G20 and G21 within the GGAG recognition motif. Combined mutation frequencies at positions G20 and G21 are 2.0% and 1.6%, respectively, with adenines clearly dominating the mutant nucleotide identities (Fig. 4B, top panel). Additionally, mutations at U8 and G23 were more prevalent in the peptide-modified RNA sample compared with controls. However, their overall mutation frequencies were lower than that observed at G20 and G21 (0.53% and 0.29%, respectively).

Because they fall within the well-annotated recognition motif for LIN28A, G20 and G21 participate in numerous interactions with the LIN28 ZKD (Fig. 5A, B). However, MS analysis of crosslinked preEM-let-7f: LIN28 complexes did not detect crosslink sites within the ZKD. To determine whether crosslinking at those sites is feasible in isolation, we sought to crosslink the LIN28 ZKD in vitro, in the absence of the LIN28 CSD. To that end, a well characterized, recombinant LIN28 ZKD construct (residues 120–181) (See ref. 39) was overexpressed and purified from E. coli. Complexes of ZKD with a corresponding fragment of pre-let-7 RNA (sequence: UAGGAGAU) were formed in vitro and crosslinked as described previously. SDS-PAGE analysis confirmed that the ZKD alone was capable of crosslinking with a corresponding RNA fragment despite low binding affinity (K_D > 7.2 uM, unquantified) (Fig. 5C, D).

**Discussion**

CLIP methods have identified thousands of putative mRNA targets of LIN28, with a significant number of binding sites in both translated and untranslated regions of the transcriptome. While LIN28 unambiguously inhibits processing of the let-7 precursors it targets, reports indicate that LIN28 can both enhance and suppress translation of subsets of RNAs with numerous cellular consequences, prompting questions about the specificity and pervasiveness of interactions, as well as concerns over reproducibility and experimental variation. The nature of LIN28 recognition of RNA likely further complicates target identification efforts as structural and biochemical studies have revealed that LIN28 recognizes let-7 precursors through a bipartite interaction mediated by two binding domains with distinct recognition characteristics.

To identify precise crosslink sites in a model recombinant LIN28A protein-RNA complex, we first used LC-MS and MS/MS to analyze covalent RNA-peptide heteroconjugates. Of the 11 ssRNA bases that mediate the LIN28A-ΔΔpreEM-let-7f crosslinks...
complex, our MS analysis identified only a single crosslink site. The covalent crosslink between Phe55 of the LIN28A CSD and a terminal loop uridine of preEM-let-7f, corresponds to a tight interfacial contact observable in the crystal structure of the complex.\textsuperscript{18} We separately confirmed this assignment using a novel, single-nucleotide isotope labeling technique.\textsuperscript{38} Aside from the single crosslink of uridine to Phe55, we were unable to identify directly any other site of modification by mass. Alternatively, our CIMS analysis of human and mouse CLIP data sets revealed an enrichment of mutations occurring at guanines, presumably indicative of crosslinking sites. Furthermore, sequencing analysis of our \textit{in vitro} crosslinked peptide modified preEM-let-7 determined an enrichment of mutations at guanines (specifically at those within the GGAG motif), and \textit{in vitro} crosslinking experiments confirmed that the ZKD can crosslink independently of the CSD. However, CIMS analyses did not determine a significant frequency of mutations at the uridine crosslink site detected by mass analysis.

Overall, our data indicate that the two analytical methods identify different crosslinking sites within the same UV crosslinked protein-RNA complex. Consistent with this finding, a recent report (see ref. \textsuperscript{41}) demonstrated that in a systematic MS/MS analysis of 124 RBPs crosslinked \textit{in vivo}, MS-detectable crosslinking occurred almost exclusively at uridines. In one data set, 89% of crosslinking events were confirmed at uridines, and at least one uridine was present in crosslinked di- and tri-nucleotides in the remaining 11% of cases, although LIN28 was not included in that report. Furthermore, they noted that protein-RNA crosslinks are typically few in number (\textasciitilde{}1–3 crosslinks per complex) and are frequently mediated by phenylalanine residues (24% of identified amino acid crosslink sites). On the other hand, the only previous report to feature CIMS profiles of LIN28 CLIP data sets (see ref. \textsuperscript{24}) identified primarily guanine points of mutation. Furthermore, the cross-link site mutation profile determined specifically for pre-let-7f showed mutations primarily at guanines within the GGAG motif and no mutations at the uridine site that we identified \textit{in vitro}, or at any uridine within the terminal loop.

Collectively, the results of our case study of existing LIN28 binding site data and our biophysical investigation of LIN28A crosslinking to a let-7 precursor suggest that the two highest resolution crosslink site detection methods may benefit from orthogonal, complementary analyses to gain complete crosslink site identification data. Given that precise identification of protein-RNA interactions on a global scale provides an invaluable tool for mechanistic and functional studies of various cellular processes, further development of comprehensive analysis methods is needed to identify, reproducibly and with high confidence, the precise binding sites between diverse RBPs and their RNA targets. One potential opportunity might be found in the advancement of CLIP methods that examine crosslinking between individual domains of proteins and their RNA targets, such as iDo-PAR-CLIP (individual domain PAR-CLIP),\textsuperscript{26} with the addition of a higher precision MS analysis step. This version of PAR-CLIP relies on a proteolytic cleavage site between two binding domains of an RBP to facilitate single domain isolation and domain specific target sequencing following UV irradiation.\textsuperscript{26} Furthermore, unambiguous and comprehensive structural data generated from key techniques such as X-ray crystallography and nuclear magnetic resonance (NMR) may be used to further scrutinize or validate CLIP determined binding sites.

Material and methods

Expression and purification of LIN28A proteins

All recombinant LIN28A protein constructs were overexpressed from pET21a or pETDuet expression vectors. BL21 Rosetta cell colonies transformed with construct plasmids were used to inoculate 100ml LB starter cultures and incubated overnight (\textasciitilde{}18–20 hrs) in a shaker incubator at 37°C. The next day, 10 ml of starter culture was used as inoculation for every 1 L of expression culture (typically 2–4 L). Expression cultures were incubated at 37°C until reaching O.D. of 0.6–0.8, at which point cultures were induced with 0.5 mM IPTG. Following induction, cultures were incubated overnight at 18°C with shaking and harvested the next day via centrifugation. Proteins were purified via Ni\textsuperscript{2+} affinity, cationic exchange and size exclusion chromatography, as described previously.\textsuperscript{18}

Electrophoretic mobility shift assays (EMSA) and SDS-PAGE

For binding analyses, LIN28A was serially diluted into a low-salt binding buffer (20 mM Bis-Tris, pH 7, 100 mM NaCl, 50 \(\mu\)M ZnCl\textsubscript{2}, 5% glycerol and 5 mM DTT) supplemented with yeast tRNA (to a final concentration of 1 mg/ml) and the RNase inhibitor Ribolock (ThermoFisher Scientific, CAT: EO0381). Precursor let-7 RNA probes were synthesized from IDT and labeled with \(^{32}\text{P}\) via the T4 Polynucleotide Kinase (New England BioLabs, CT: M0201S). \(^{32}\text{P}\)-labeled RNA substrates (< 1 nM) were incubated with LIN28A protein dilutions of increasing concentration for \textasciitilde{}30 minutes at RT before samples were run on a 10% native gel. Gels were vacuum dried and pressed to radiolabel sensitive film overnight. Films were imaged the following day using scanning phosphorimager. For SDS-PAGE analyses, complexes of LIN28A proteins with pre-let-7f RNAs were exposed to UV irradiation and pre- and post-crosslinked samples were compared on SDS-PAGE gels and stained with Coomassie Blue.

UV crosslinking and hydrolysis of LIN28A and let-7 complexes

Purified complexes of LIN28A proteins and preE-let-7f targets were buffer exchanged into crosslinking buffer (20 mM Bis-Tris, pH 7, 100 mM NaCl, 50 \(\mu\)M ZnCl\textsubscript{2}, and 1 mM DTT), and UV crosslinking was completed by irradiating samples three times at 300 mJ/cm\textsuperscript{2} at 254 nm in a Stratalinker 1800. Crosslinked complexes were separated from non-crosslinked complexes via denaturing urea gel. Gels were stained with ethidium bromide and imaged. Crosslinked complex RNA bands (indicated by the increase in size over the free RNA bands) were excised and RNA was eluted from gel via electro-elution into a dialysis bag, 3 \times 30 mins at 100V. Eluted, crosslinked complexes were concentrated and buffer exchanged into 8 M urea, 50 mM bis-Tris, pH 7. Trypsin/Lys-C (Promega, CT: 1762 E. RANSEY ET AL.
V5072) was added to a final ratio of 1:25 (w/w). Digestions were incubated at 37°C for 3 hrs with shaking. Samples were diluted to lower the urea concentration to < 1 M and digestions continued overnight at 37°C.

Modified RNA was enriched from digested samples via anion exchange chromatography. Specifically, samples were loaded onto a DEAE column (GE Healthcare) with low-salt buffer (20 mM Bis-Tris, pH 6, 10% glycerol and 5 mM DTT) and eluted with a high salt fast gradient (20 mM Bis-Tris, pH 6, 10% glycerol and 5 mM DTT, 2 M NaCl). Chromatography fractions containing RNA were pooled, flash frozen, and lyophilized. Dried samples were resuspended in LC-MS grade water and quantified by their optical density at 260 nm.

**LC-MS**

All samples were separated on an Agilent 1200 HPLC coupled to a solvent degasser, auto sampler, diode array detector, and column oven (Agilent Technologies). Separations were performed using two solvent systems. Native and trypic peptide-modified, full-length RNA samples were separated on a 100 mm × 1 mm i.d. Xbridge C18 column with a particle size of 3.5 μm (Waters). The solvent system was based on a previously published reverse phase ion pairing LC-MS method, and we used 200 mM HFIP with 1.25 mM trimethylamine at pH 7.0 in mobile phase A and methanol in mobile phase B. The column was heated to 60.0°C and the flow rate was 100 μL/min. Injection volumes were 10 – 25 μL. Mobile phase B was increased from 5% to 15% from 0 – 20 minutes and then from 15% to 60% over an additional 20 minutes of run time. Absorbance was monitored at 260 nm with a reference wavelength at 380 nm and a 2 s response time. Tryptic, crosslinked peptides digested in formic acid were separated on a 150 mm × 1.0 mm i.d., micro bore rapid resolution SB-C18 column with a particle size of 3.5 μm (Agilent Technologies). Mobile phase A was water with 0.1% FA and mobile phase B was acetonitrile with 0.1% FA. The column was heated to 40.0°C, and the flow rate was 150 μL/min. Injection volumes were 5 – 20 μL. Mobile phase B was increased from 5% to 45% over 0 – 20 minutes.

All samples were analyzed on an Agilent G6520A accurate-mass QTOF coupled to the LC system described above, operating in extended dynamic range mode. The system was calibrated on the same day of analysis and reference masses were continuously infused for online mass correction. Enriched RNA oligonucleotides crosslinked to tryptic peptides were separated using the HFIP/TEA solvent system and analyzed in negative ion mode from 239 – 3200 m/z with a scan rate of 1 spectrum/s using the following settings: drying gas flow, 8 L/min; drying gas temperature, 325°C; nebulizer pressure, 35 psig; capillary voltage, 4500 V; fragmentor, 175 V; and skimmer: 65 V. In targeted MS/MS mode, ions were fragmented using collision induced dissociation with nitrogen gas, collision potentials of 10 – 30 V and an isolation width of ~4 amu.

**LC-MS data analysis and crosslink site identification**

Tryptic peptides crosslinked to mono- or dinucleotides were first identified based on MS1 data using a database composed of predicted trypptic peptides with up to two missed cleavages crosslinked to all possible mono- and dinucleotides using Agilent’s Find by Formula algorithm in the MassHunter software package. The analysis allowed for the loss of water so we could identify both mass neutral crosslinks and those leading to loss of water. The peptides identified in survey spectra with covalent UMP, di-, and trinucleotide modifications were then analyzed in targeted MS/MS experiments to validate the peptide sequence and characterize the species. The targeted MS/MS spectra for the heteroconjugate MGFGLMSMTR-UMP were manually compared with theoretical product ion spectra in which the UMP was attached to different amino acids in the peptide as to determine the position of the crosslink.

**Workflow for CIMS analysis**

CLIP data sets for mouse (ref. 24) and human (ref. 23) LIN28 were obtained through Gene Expression Omnibus (GEO), with accessions SRR458758 (monoclonal 353L33G), SRR458760 (polyclonal), SRR531464 (LIN28A) and SRR531465 (LIN28V5). The WormBig and WormSmall data sets from ref. 27 were transferred via personal communication. The SRA files were downloaded and converted to FASTQ files using SRA Toolkit 2.5.2. FastQC determined the sequence quality and presence of adapters. Bowtie2 indexes were created for hg19, mm10 and ce10 for human (H. sapiens), mouse (M. musculus), and worm (C. elegans), respectively. Bowtie2 was run with the parameters -p 18 -N 1. Samtools was used to sort and convert mappings into SAM files. The SAM files were then read into R and the following filters applied: low quality reads were removed (if for any nucleotide the quality is below D then the whole read is removed) and all exact sequence duplicates were collapsed to eliminate the potential for PCR duplicates. For each mutation type (substitutions/deletions), distributions were derived separately. For substitutions, only reads with a single substitution were considered. Substitution positions were identified based on the SAM files, fields for tag MD type Z. For deletions, only reads with a single deletion were considered. The positions of deletions were identified based on CIGAR string based on the SAM files. Stringent conditions for the number of substitutions and deletions (1 per read) led to filtering out reads with adaptor sequences. Further analyses of obtained distributions were performed with ggplot2 package, and were visualized with ggplot2 package for R.
Peptide-modified preEM-let-7f sequencing library generation and analysis

Purified, lyophilized peptide-modified preEM-let-7f was resuspended in LC-MS grade water and quantified via optical density measurement at 260 nm. The QiaSeq miRNA library construction kit (Qiagen, Cat#331502) was used to generate a library from ~100ng of peptide-modified RNA using the standard protocol, incorporating 16 cycles of PCR. For quality assurance, Tape Station High Sensitivity tape is run on the Tape Station 2200 from Agilent. Additionally, a qPCR run was completed using the Kapa Library Quantification Kit (Kapa, KK4824) with a standard 0 for normalization. Generated libraries were pooled together in equimolar proportions and were denatured per standard Illumina protocol. Sequencing was performed on an Illumina MiSeq with Nano protocol at 6 pM with 50% PhiX for a read length of PE 75 molar proportions and were denatured per standard Illumina protocol. Sequencing was performed on an Illumina MiSeq with Nano protocol at 6 pM with 50% PhiX for a read length of PE 75 and depth of up to 1 million reads.

Sequencing analysis of peptide modified preEM-let-7f library

Reads were trimmed with cutadapt to remove the 3’ adaptor. Bowtie2 was used to create the index of the reference sequence. Then, they were aligned with bowtie2 with parameters: -D 20 -R 3 -N 0 -L 10 -i S,1,0,5 to allow short alignments. Samtools was used to sort and index the alignment file, and sambamba-depth was used to calculate the coverage along the reference sequence. Substitution ratios were calculated in R dplyr and results were visualized with ggplot2.

Conflict of interest statement

The authors declare that there are no conflicts of interest

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Abbreviated title

Comparative analysis of LIN28-RNA UV-induced crosslinks

ORCID

Elizabeth Ransey https://orcid.org/0000-0002-8118-466X
Anders Bjorkholm https://orcid.org/0000-0003-0611-4175
Victor S. Lelyveld https://orcid.org/0000-0002-3890-0288
Przemyslaw Bipecz https://orcid.org/0000-0001-8423-1823
Lorena Pantano https://orcid.org/0000-0002-3859-3249

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