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An RNA-binding Protein, Lin28, Recognizes and Remodels G-quartets in the MicroRNAs (miRNAs) and mRNAs It Regulates*

Elizabeth O’Day†‡, Minh T. N. Le§, Shunsuke Imai¶, Shen Mynn Tan†, Rory Kirchner¶, Haribabu Arthanari‡, Oliver Hofmann*, Gerhard Wagner†1, and Judy Lieberman‡2

From the †Department of Biological Chemistry & Molecular Pharmacology, Harvard Medical School, Boston, Massachusetts 02115, the §Cellular and Molecular Medicine Program, Boston Children’s Hospital, Boston, Massachusetts 02115 and ‡Department of Biostatistics, Harvard School of Public Health, Boston, Massachusetts 02115

Background: Lin28 is an evolutionary conserved RNA-binding protein that regulates miRNAs and mRNAs; Lin28 overexpression is linked to poor prognosis in cancer.

Results: Lin28 binds to guanine-rich miRNAs and mRNAs that contain G-quartet structural features and remodels these structures.

Conclusion: Lin28 recognition of G-quartets is a unifying feature of Lin28-regulated RNAs.

Significance: Small molecules that bind to G-quartets might be useful inhibitors of Lin28 function.

Lin28 is an evolutionary conserved RNA-binding protein that inhibits processing of pre‐let‐7 microRNAs (miRNAs) and regulates translation of mRNAs that control developmental timing, pluripotency, metabolism, and tumorigenesis. The RNA features that mediate Lin28 binding to the terminal loops of let-7 pre‐miRNAs and to Lin28‐responsive elements (LREs) in mRNAs are not well defined. Here we show that Lin28 target datasets are enriched for RNA sequences predicted to contain stable planar structures of 4 guanines known as G‐quartets (G4s). The imino NMR spectra of pre‐let‐7 loops and LREs contain resonances characteristic of G4 hydrogen bonds. These sequences bind to a G4‐binding fluorescent dye, N‐methylmesoporphyrin IX (NMM). Mutations and truncations in the RNA sequence that prevent G4 formation also prevent Lin28 binding. The addition of Lin28 to a pre‐let‐7 loop or an LRE reduces G4 resonance intensity and NMM binding, suggesting that Lin28 may function to remodel G4s. Further, we show that NMM inhibits Lin28 binding. Incubation of a human embryonal carcinoma cell line with NMM reduces its stem cell traits. In particular it increases mature let‐7 levels, decreases OCT4, HMGAl, CCNB1, CDK4, and Lin28A protein, decreases sphere formation, and inhibits colony formation. Our results suggest a previously unknown structural feature of Lin28 targets and a new strategy for manipulating Lin28 function.

Lin28 regulates networks of RNAs that control developmental timing (1–3), pluripotency (4, 5), metabolism (6, 7), and tumorigenesis (8–10). Lin28 was first identified as a heterochronic gene coordinating the “larva-to-adult” switch in Caenorhabditis elegans (1, 2, 11). All bilaterian animals possess a Lin28 homologue. It is abundantly expressed in embryos and stem cells, but absent in most terminally differentiated cells with the exception of skeletal and cardiac muscle (12, 13). In embryonic stem cells, Lin28 contributes to pluripotency by promoting self-renewal and blocking differentiation (14–17). It is part of a small unique set of proteins that include OCT4, SOX2, and Nanog that can reprogram terminally differentiated fibroblasts to induced pluripotent stem cells that resemble embryonic stem cells (18). Lin28 is the only protein that is not a transcription factor identified thus far to assist in reprogramming, suggesting that Lin28‐targeted RNAs regulate stem cell properties.

Lin28 overexpression is associated with many human cancers and is linked to poor prognosis (16). Approximately 15% of primary tumors and human cancer cell lines have increased Lin28 expression (8, 9, 16, 19). Elevated Lin28 is associated with aggressiveness of germ cell tumors, hepatocellular carcinoma, and cancers of the breast, colon, thyroid, esophagus, ovaries, prostate, and head and neck (8, 9, 20–22). Ectopic Lin28 expression facilitates cellular transformation, and Lin28‐overexpressing cells can form tumors in mice (9). Expression of the Lin28 target let‐7 inhibits tumor formation and metastasis (23–25). Uncovering the mechanism by which Lin28 selects RNA targets and regulates them is critical to understanding the function of Lin28.

The Lin28 protein contains three nucleic acid-binding domains, a cold‐shock domain (CSD)3 and two zinc finger (ZnF) motifs. Lin28 is the only known animal protein that contains both a CSD and ZnFs. Lin28 binds to the loops of pre‐let‐7
miRNAs to block processing by Dicer (17, 26, 27). The loop sequences are diverse in size and sequence. Upon binding to let-7 pre-miRNAs, Lin28 recruits ZCCHC11, also known as TUTase4 (TUT4), a uridylyl transferase, which adds a 3’-uridine tail to pre-let-7s (27–30). Terminal uridylation promotes pre-let-7 degradation.

Although much of Lin28 research has focused on its role in regulating let-7 biogenesis, miRNAs constitute the overwhelming majority of RNAs bound to Lin28 in cells (31). There is no agreement in the literature about common features of Lin28-binding sites. In vitro binding assays suggested that specific guanine (26, 32, 33), adenosine (26), uracil (34), and cytosine (35) nucleotides affect Lin28 binding. The crystal structures of Lin28 bound to pre-let-7 loop oligonucleotides suggested that the CSD and ZnF domains form critical contacts with guanine and adenine bases in distinct G-rich regions in the loops of pre-let-7s (33, 36, 37). In particular the ZnF domain was proposed to bind to a GGAG motif in the loop sequence of pre-let-7 miRNAs. However, the NMR structure of isolated ZnFs complexed with AGGAGAU suggested that an exact GGAG sequence is not required (36). The Lin28 CSD appears to have little sequence specificity because it forms crystals with d(T)6 and r(U)6 (37). Sequencing of Lin28-bound mRNAs suggested short Lin28 recognition elements. Such sequences are diverse in size and sequence. Upon binding to miRNAs to block processing by Dicer (17, 26, 27). The loop structures of the pre-let-7s and their loops, which contain the putative Lin28 consensus sequences AAGNNG, AAGNG(N), or (N)UGUG(N), where N is any nucleotide (31, 38). Molecular modeling instead suggested that Lin28 recognizes a critical “A bulge” flanked by two G:C bonds in an extended double-stranded region of the LREs of mRNAs (39). A single A to U mutation in OCT4, HMGA1, and RPS19 mRNAs inhibited the ability of Lin28 to stimulate their translation (39). However, the “A bulge” is not present in pre-let-7 miRNAs.

These structural, sequencing, and modeling data have suggested short Lin28 recognition elements. Such sequences are abundant in miRNAs throughout the transcriptome and hence are unlikely to uniquely define Lin28 binding specificity. None of these motifs are useful for predicting LREs of Lin28-regulated miRNAs. Thus, the key RNA features that determine Lin28 binding remain unclear.

We hypothesized that Lin28-regulated miRNAs and miRNAs share common structural features. We observed that Lin28 targets have an abundance of guanine and guanine repeats. Because guanine rich nucleic acids can form stable planar structures called G-quartets (G4s) by G:G:G:G hydrogen bonding, we hypothesized that Lin28 might recognize structured RNAs that contain G4s. To address this hypothesis, we used bioinformatics, NMR spectroscopy, a fluorescent dye that specifically binds to G4s, and gel shift assays to analyze the structural features of the pre-let-7s and their loops, which contain the 3’-GGAG motif; the LREs of DSG2, HSPA5, and SUN1, which contain the putative Lin28 consensus sequences AAGNNG, AAGNG(N), and (N)UGUG(N), respectively; and the LRE of OCT4, which contains the proposed “A bulge.” We provide evidence that a G-quartet structure is a unifying feature of Lin28 RNA targets. Further, we demonstrate that Lin28 may function to unwind G4s and that G4-stabilizing agents represent a novel class of Lin28 inhibitors.

Experimental Procedures

**RNA Sequences**—Sequences of RNA used in this study are as follows: pre-let-7d, 5’-CCUAGAGGAAGGUAGGUUGCG- AUAGUUUAGGGCAGAUUUGCACCACAGGAGGUAACUAUAGGACCUGGCUCGUUUUCUUAGG3’; let-7d-loop, 5’-UUAGGGCCAGGAUUUGCCACAAAGGAGGUAACUAUAGGACCUGCCGUUUACCCCUAGG3’; pre-let-7g, 5’-AGGCGUGAGAUAGUUGUUAAGGUUGCGGAGGGAUGGUGGCACCAACAGGAGGUAACUAUAGGACCUGCCGUUUACCCCUAGG3’; let-7g-loop, 5’-GAGGCGUGAUAUACCAACCCGGUACAGGAGG3’; DSG2 LRE, 5’-GGUCUGCAUAUUGGAGCCAGG3’; HSPA5 LRE, 5’-GGCGUGAGGAUAAAGCGGUAGG- AGCCUUGCUUU-3’; SUN1 LRE, 5’-UAUCCAGAAGGAAGGUGAGGAAG-3’; OCT4 LRE, 5’-GCAGAAGAGGACCCUUGGGGAUAAACACAGCCGGAGUGGGGUCCACCCUGGGGUUCUAUUUGGAGAUUAUCGCAAACGCACAUUGCCC-3’; miR-21, 5’-GAUUGUAGACGUUUAGUAUCUAGACACAC-3’.

**RNA Synthesis and Purification**—For NMR studies, template DNA containing the T7 promoter, sequence of interest, and restriction enzyme cleavage site for linearization (Smal or DraI) were generated by PCR and inserted into the pUC19 plasmid after EcoRI and BamHI digestions. After linearizing the plasmid with Smal or DraI, the templates were amplified by PCR, extracted with phenol-chloroform, and precipitated with ethanol.

RNAs were synthesized from this template by in vitro transcription using T7 RNA polymerase, in 10–20-ml reactions. After denaturation at 95 °C for 2 min, the RNAs were precipitated with ethanol and purified by electrophoresis on urea-containing polyacrylamide denaturing gels. Note that these RNAs have 1–3 additional guanosines at their 5’ termini for efficient transcription, and one additional uracil (HSPA5) or 1–3 additional cytosines (the others) at their 3’ termini for linearization of the template by DraI or Smal, respectively.

**NMR Spectroscopy**—NMR spectra were acquired on Bruker 500- or 600-MHz spectrometers equipped with cryoprobes. One-dimensional 1H NMR spectra were acquired after incubating 100–500 μM RNA in 10 mM sodium phosphate buffer, pH 6.4, 100 mM KCl, 4 mM MgCl2 in 90% H2O/10% D2O. Samples were equilibrated at 5 °C. Data were accumulated with 1,024 transients over a frequency width of 10.33 kHz with 32,768 data points. Spectra were processed using MestReNova.

**N-Methyl Mesoporphyrin IX (NMM) Fluorescence Assay**—NMM was obtained from Frontier Scientific (Logan, UT). Fluorescence assays were performed in 30 μl of 10 mM sodium phosphate buffer, pH 6.4, 100 mM KCl, 4 mM MgCl2, and 5 μM NMM. The RNA concentration ranged from 0 to 100 μM. All fluorescence experiments were performed using a FlexStation III plate reader with excitation and emission wavelengths of 399 nm and 614 nm, respectively. Fluorescence measurements were repeated three times for each sample, and the intensities were averaged and corrected by running a buffer control without RNA before each series of experiments. Fluorescence intensities were normalized to the maximum intensity of UG4U. Results shown are the average of 3–5 independent replicates.
Lin28 Recognizes and Remodels G-quartets

Lin28 Expression and Purification—The cDNA for mouse Lin28A were obtained from Addgene (Cambridge, MA). The region containing the CSD and ZnFs (residues 36–186) was PCR amplified and cloned into the bacterial expression vector pET15b, adding a C-terminal His6 tag. Lin28-His6 was overexpressed in Escherichia coli Rosetta (DE3) cells (Millipore). Cells were grown in Luria Broth (LB) at 25 °C to an A600 of 0.9 and induced with 1.0 mM isopropyl β-D-1-thiogalactopyranoside. After 3 h, cells were harvested by centrifugation. Cells were lysed by sonication in 50 mM Tris-Cl, pH 7.6, 100 mM KCl, 5 mM β-mercaptoethanol, 0.1 mM ZnCl2, and 10 units of Benzonase nuclease (Sigma). Lin28-His6 was purified by nickel affinity chromatography followed by size-exclusion chromatography on a Superdex 75 prep-grade column in 50 mM Tris-Cl, pH 6.4, 200 mM NaCl, 5 mM β-mercaptoethanol. All purification steps were carried out at 4 °C.

Electrophoretic Mobility Shift Analysis—EMSA experiments were conducted using 2 × 106 cpm 32P-5′-end-labeled RNA probes. Binding buffer contained 50 mM Tris (pH 7.6), 100 mM KCl, 5% glycerol, 40 units of RNaseOUT (Invitrogen), and 10 mM β-mercaptoethanol. Binding reactions were conducted in 20 μl with increasing amounts of recombinant Lin28-His6 (0, 0.15, 0.3, 0.6, 1.25, 2.5, 5.0, 10 μM). Bound complexes were resolved on native 5% polyacrylamide gels and visualized by autoradiography.

Unwinding Assay—Unwinding assays were carried out with constant amounts of let-7g loop or HSPA5 LRE (50 μM) in 10 mM sodium phosphate buffer, pH 6.4, 100 mM KCl, 4 mM MgCl2, and 5 μM NMM to which recombinant Lin28 or BSA protein was added. The protein concentration ranged from 0 to 6.25 μM. After the addition of protein, samples were allowed to equilibrate at least 4 h at 4 °C. Fluorescence intensities were recorded using a FlexStation III plate reader, as in the NMM fluorescence assay. Protein only samples were recorded in 10-cm plates in serum-containing medium. Medium was replaced every 3 days. After 8–14 days, cells were fixed in methanol (∼20 °C) and stained with crystal violet.

Luciferase Reporter Assay—Human NCCIT cells were transfected alone, with 5 nM wild-type pre-let-7g (5′-AGGCCUGAGGUAGUAGUCAUGUUGAGGGUCUAUGUAUACCACCCGGUACAGGAGAUACUGACCACUGGCCUUGCCA-3′), or 5 μM mutant pre-let-7g (5′-AGGCCUGAGGUAGUAGUCAUGUUGAGGGUCUAUGUAUACCACCCGGUACAAAAGAUGACUGACCACUGGCCUUGCCA-3′) and 25 ng of psiCHECK3 (Promega) vector containing two copies of the let-7 binding site cloned into the multiple cloning site (NotI and Xhol) of Renilla luciferase. After 48 h, cells were incubated with 100 μM NMM for 3 h and luciferase activities were measured using the Dual-Luciferase assay system (Promega). Data were normalized to firefly luciferase.

Results

Lin28-regulated Gene Sequences Are Enriched for G-quartet Motifs—We examined the sequences of the 12 let-7g loops and of all 15 Lin28 targets with an experimentally defined LRE (Table 1). Many of the LREs are enriched for Gs and have multiple stretches of 2–6 consecutive guanines. Those LREs that do not have an excess of Gs are mostly longer (200–3270 nucleotides) and may not correspond to the minimal LRE sequence. G-rich nucleic acids can form structures containing G4s (Fig. 1A). Stacked G4s adopt unique structures known as G-quadruplexes that form diverse structures (Fig. 1B) depending on the number of stacked G4s, strand orientation, loop structures, and whether G4s are formed via inter- or intramolecular interactions (40–42). To determine whether Lin28 RNA targets might contain G4s, we first utilized the Quadruplex G-Rich Sequence (QGRS) Mapper, which predicts G4s in nucleic acids by identifying potential G-quartet sequences that match the following motif: GₙNₓGₓ₂Gₓₙ₋₂Gₓₙ₋₂Gₓₙ₋₂Gₓₙ₋₂, where nx is the number of G-repeats and y₁,y₂, and y₃ are the lengths of the gaps (43). The program generates a G-score such that sequences with higher scores are more likely to form G4s. QGRS was applied to the 50 most enriched mRNAs immunoprecipitated with Lin28 antibody in human embryonic stem cells (5) and the 50 most enriched Lin28-bound mRNAs identified by photoactivatable ribonucleoside-enhanced cross-linking and immunoprecipitation (PAR-CLIP) in FLAG/HA-Lin28-overexpressing HEK293 cells (34) (Fig. 1C). (We did not apply this analysis to putative Lin28 mRNA targets identified by two other studies that gave large numbers of Lin28-interacting mRNAs (6,000 and 13,000) without any ranking (31, 38)). The G scores of Lin28-bound RNAs were compared with the scores of the 50 mRNA most...
enriched in binding to FMRP (fragile X mental retardation protein-1), which is known to contain G4s (44–46), as positive control, or HuR, which binds AU-rich mRNAs (47), as negative control. These datasets were compared with the G-scores of 50 randomly chosen mRNAs. As expected, FMRP-binding mRNAs had significantly higher G-scores than random mRNAs (p = 0.017), and the HuR mRNAs had significantly lower G-scores than random mRNAs (p < 0.0001) (Fig. 1C). The top 50 Lin28-bound mRNAs in the immunoprecipitation and PAR-CLIP datasets had significantly higher G-scores than the random mRNAs (p < 0.0001 and p = 0.017, respectively). The increased G-scores of Lin28-bound mRNAs suggests that a common feature might be the ability to form G4s.

Lin28-regulated miRNAs and mRNAs Contain G-quartet Features—By monitoring imino protons, NMR can detect the hydrogen-bonding network that governs nucleic acid structure. Labile imino proton resonances are only observed when they are protected from solvent exchange by hydrogen bonding. In Watson-Crick base pairing, A:U hydrogen bonds have a chemical shift at ~13 ppm, whereas G:C hydrogen bonds are closer to 12.7 ppm (48). G:U wobbles and other non-canonical hydrogen bonds occur upfield of 12.7 ppm (48, 49). The unusual hydrogen-bonding pattern of G4s (Fig. 1A) gives rise to characteristic imino resonances at ~10–12.2 ppm (48, 49). Peaks within this region are highly suggestive of G4 formation. We recorded one-dimensional proton NMR spectra for Lin28-binding RNAs (pre-let-7d, the let-7d loop, pre-let-7g, the let-7g loop, and the LREs of DSG2, SUN1, HSPA5, and OCT4). All of the RNAs had imino resonances indicating the presence of non-canonical hydrogen bonds (Fig. 2A). In addition to putative A:U and G:C bonds, each Lin28 RNA target had resonances in the non-canonical hydrogen bond region from ~10.0 to 12.2 ppm. Imino resonances in this region suggest that pre-let-7d, the let-7d loop, pre-let-7g, the let-7g loop, and LREs of DSG2, SUN1, HSPA5, and OCT4 might contain G4 features. We attempted to use multidimensional NMR methods including NOESY experiments to unambiguously map the hydrogen bond connectivity of these Lin28-binding RNAs. However, from PAGE we observed that these RNAs oligomerize and/or dynamically change structure in solution, making additional NMR analysis difficult to interpret. Thus, although the imino resonance pattern suggests the presence of G4s, we were unable to obtain definitive structural evidence.

To gain further insight as to whether Lin28 RNAs contain G4s, we made use of NMM, a dye that binds selectively to G4s (52–55). When it binds to G4s, NMM is excited at 399 nm and fluoresces at 614 nm (55). We therefore compared NMM fluorescence of these Lin28-binding RNAs (pre-let-d, the let-7d loop, pre-let-7g, the let-7g loop, and the LREs of DSG2, SUN1, HSPA5, and OCT4) with that of RNAs that lack a G4 structure. Strong NMM fluorescence was measured for the known G4 RNA UG4U (UGGGGU) and for all of the Lin28-binding RNAs measured, but little to no fluorescence was detected when NMM was mixed with yeast-tRNA, poly(U), or the loop of miR-21 (Fig. 2B). Taken together with the one-dimensional NMR results, NMM fluorescence suggests that Lin28-binding miRNAs and mRNAs form G4s.

**RNA Mutations That Disrupt Lin28 Binding Disrupt NMM Fluorescence**—To probe whether Lin28 binding to RNA correlates with G4 formation, we generated RNAs that were truncated or contained mutations that might affect G4 formation. For the let-7g loop, we mutated C18C19 to U18U19 (C-Mt), G27G28 to A27A28 (G-Mt1), and G4G5 and G27G28 to A4A5 and A27A28 (G-Mt2). We also synthesized an HSPA5 LRE truncation by removing 9 nucleotides from the 5’-end containing the first two GG repeats (HSPA5-tr). We tested the ability of each mutant to bind Lin28 by EMSA and to bind to NMM by fluorescence assay (Fig. 3). When compared with the wild-type let-7g loop, the C-Mts had increased affinity for Lin28 as suggested by the relative intensity of the free let-7g loop with the gel-shifted band and increased NMM fluorescence. G-Mt1 had reduced Lin28 binding and weaker NMM fluorescence. G-Mt2 did not bind to Lin28 and showed no NMM fluorescence. HSPA5-tr did not cause a Lin28 gel shift or NMM fluorescence.
suggesting that G-repeats are critical for Lin28 binding and that Lin28 binding correlates with NMM fluorescence. These findings suggest that Lin28 and NMM recognize a shared structural feature of RNA. Because NMM is only known to bind G4s, Lin28 likely recognizes a G4 structure in its targets.

Lin28 Binding Disrupts the Ability of Lin28 RNA Targets to Interact with N-Methyl Mesoporphyrin IX—Upon binding, Lin28 induces a conformational change in its RNA targets (32, 36, 37, 56). We hypothesized that the Lin28-induced RNA rearrangement might remodel the RNA G4 structure. To test this idea, we monitored changes to the imino protons of the let-7g loop and the HSPA5 LRE upon the addition of Lin28 (Fig. 4A). When saturating amounts of Lin28 were incubated with these RNAs, the intensity of the iminos decreased and the G4-like imino signals were greatly reduced. This suggests that Lin28 directly binds to these target RNAs and that binding disrupts the hydrogen-bonding network. To further examine the ability of Lin28 to unwind G4s, we performed a modified NMM assay in which the let-7g-loop or HSPA5 LRE was incubated with a constant concentration of NMM and increasing amounts of rLin28A or BSA (Fig. 4B). The let-7g-loop and HSPA5 LRE exhibited strong NMM fluorescence on their own, but when Lin28 was added, NMM fluorescence was lost. The addition of BSA had no effect. Thus, in the presence of Lin28, the RNA targets are no longer able to interact with NMM. These results together with the NMR data suggest that Lin28 binding remodels the G4s.

NMM Inhibits Lin28—If remodeling G4s is crucial to Lin28 function, the G4 feature of Lin28-binding RNAs could represent a novel site to manipulate Lin28 activity. By binding to G4s
FIGURE 2. **Lin28 RNA targets contain G4 features.** A, $^1$H imino proton spectra for Lin28 targets contain resonances indicative of non-canonical hydrogen bonds similar to G4s (boxed region). B, Lin28 targets bind to NMM, a G4-specific fluorescent dye, but poly(U), yeast tRNA, and the loop of miR-21 do not. Error bars indicate ± S.D.
within gene promoters and within regulatory sites of mRNAs, G4-intercalating agents, like NMM, inhibit transcription and translation (57–59). We performed a competition EMSA to examine whether NMM inhibits Lin28 binding (Fig. 5A). Both 32P-labeled let-7g-loop and HSPA5 LRE bound rLin28A as measured by the presence of a gel shift. However, adding increasing amounts of NMM led to a decrease in the gel shift and an increase in the signal of free RNA. Thus, NMM inhibits the ability of Lin28 to bind RNA. To our knowledge, small-molecule inhibitors of Lin28 binding have not been described before.

The miR-21 loop and HSPA5 LRE truncation (HSPA5-tr) did not bind Lin28 or NMM at the highest concentration.

We next sought to determine whether NMM could inhibit Lin28 function in cells. Human NCCIT embryonal carcinoma cells, which, like embryonic stem cells, express high levels of Lin28, were incubated with 100 μM NMM. NMM had no significant effect on cell viability (Fig. 5B). However, exposure to NMM for 3 h increased levels of mature let-7a and let-7g miRNAs, but did not affect miR-21 or miR-34a control miRNAs (Fig. 5C). This increase was specific to mature let-7 miRNAs as neither pri-let-7 nor pre-let-7 levels changed with NMM treatment. Thus, NMM alleviates the Lin28-induced block in let-7 processing.

Next we used a cell reporter assay to determine whether adding NMM increases let-7 function (Fig. 5D). NCCIT cells were transfected with a luciferase reporter containing let-7-binding sites in the 3′-UTR and either wild-type pre-let-7g or a mutant pre-let-7g, in which 4 guanines were replaced with adenosines. These mutations do not affect the mature let-7 sequence (Fig. 3D, underlined) but prevent G4 formation (Fig. 3F). The addition of both wild-type and mutant pre-let-7g decreased luciferase activity. The mutant pre-let-7g showed greater luciferase inhibition, which could result from diminished binding of Lin28 to the mutant pre-let-7g sequence (Fig. 3D). Both wild-type and mutant pre-let-7g decreased luciferase activity, but mutant pre-let-7g inhibited activity more. NMM had no effect

**FIGURE 3.** Lin28 binding correlates with NMM fluorescence. A, let-7g loop mutations and HSPA5 truncation evaluated for gel shift and NMM binding. B–E, EMSAs comparing the Lin28 interaction with wild-type and mutant let-7g loop (B–D) and full-length and truncated HSPA5 LRE (E). F, NMM fluorescence assay for wild-type and mutant let-7g loop and full-length and truncated HSPA5 LRE. Error bars indicate ± S.D.
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A

let-7g loop

HSPA5 LRE

B

Fluorescence intensity vs. Protein concentration (μM)

- G-loop + BSA
- G-loop + Lin28
- HSPA5 + BSA
- HSPA5 + Lin28
- BSA only
- Lin28 only
on mutant pre-let-7, but wild-type pre-let-7 now inhibited luciferase assay as well as mutant pre-let-7. Thus, the increase of NMM in let-7 processing led to better gene knockdown.

Lin28 also promotes translation of stem cell target genes. With NMM, we observed a decrease in the protein levels of Lin28 mRNA targets OCT4, HMGA1, CCNB1, CDK4, and Lin28A (Fig. 5E), but there was no effect on actin or tubulin protein levels. The mRNAs of the Lin28 target genes were unchanged, with the exception of MYC and Lin28 mRNA, which were significantly decreased (Fig. 5F). MYC transcription is known to be affected by G4 agents. The decrease in Lin28 mRNA suggests that NMM may also interfere with Lin28 transcription.

Because of increases in let-7 miRNAs and decreased levels of proteins associated with pluripotency, we next examined whether NMM affects stem-like properties using sphere-forming assay. The number of sphere-forming cells decreased and colony formation was blocked (Fig. 5G and H). *p value < 0.005, **p < 0.001 using a two-tailed Student’s t test. Error bars indicate ± S.D.
ing and colony-forming assays. The ability to form spheres \textit{in vitro} depends on the presence of self-renewing stem cells within a population. We counted the number of spheres embryonal carcinoma cells formed in the presence and absence of NMM. Incubation with NMM significantly reduced the number of sphere-forming cells by \( \approx 75\% \) (Fig. 5G). This suggests that NMM inhibits self-renewal. We also tested NMM in a colony formation assay, where the ability of single cells to undergo unlimited division is assessed. NMM nearly abolished the ability to form colonies (Fig. 5H). This further supports our hypothesis that NMM inhibits self-renewal and stem-like properties. Because NMM binds broadly to G4-containing RNAs and DNAs, it is unlikely that these effects are due solely to the ability of NMM to inhibit Lin28. Nonetheless, these results suggest that G4-intercalating agents can inhibit Lin28 binding to RNA and inhibit stem-like properties of cells.

Discussion

Lin28 is an evolutionarily conserved RNA-binding protein that regulates cell differentiation (12, 16). It has been unclear how Lin28 identifies its target RNAs and distinguishes them from other RNAs. Although multiple RNA sequence motifs have been suggested to influence Lin28 binding to RNA, the motifs are small and abundant in the transcriptome. Our data strongly suggest that Lin28 recognizes a common structural feature in its RNA targets; Lin28-regulated miRNAs and mRNAs share G-quartet features.

Lin28-bound mRNAs are enriched for sequences with increased potential to form G4s. The Peng dataset of Lin28 targets contains 1,200 mRNAs that were enriched at least 2.5-fold in a Lin28 immunoprecipitation when compared with a pre-immune control immunoprecipitation in human embryonic stem cells (5). No significantly enriched linear sequence was identified within this list of Lin28 targets. However, we found that the top 50 most enriched Lin28 mRNAs from this dataset are highly enriched for sequences likely to form G4s. In addition, the top 50 Lin28 mRNA targets from the Hafner dataset of Lin28-bound mRNAs identified by PAR-CLIP in FLAG/HA-Lin28-overexpressing HEK 293 cells were also predicted to have an increased likelihood to form G4s (34). The increased potential to form G4s within multiple Lin28 target datasets suggests that the G4 structure might be common to all Lin28-binding RNAs.

By one-dimensional NMR spectroscopy, we found that Lin28-binding RNAs contain imino resonances from 10 to 12.2 ppm, indicative of non-canonical hydrogen bonds (48, 51, 60). The imino resonance pattern is characteristic of the G:G:G:G hydrogen-bonding networks found in G4s (48, 51, 61). We were unable to obtain high-resolution two-dimensional NMR spec-
tra that could provide further structural support. G4-containing RNAs are prone to aggregate (51, 62–64), which may have foiled our efforts.

Lin28-binding RNAs interact with the highly specific G4 fluorescent dye, NMM (54, 55). Lin28 RNA targets, but not control RNAs such as the loop of miR-21, induced NMM fluorescence. Mutations and truncations of Lin28 RNA targets that hindered Lin28 binding also led to decreased NMM fluorescence. These results suggest that Lin28 and NMM recognize a similar structure. Because NMM selectively binds only to G4s, Lin28 RNA targets likely contain G4 features that are important for Lin28 recognition.

Recent atomic structures of Lin28 in complex with fragments of pre-let-7 loop sequences did not detect G4s (33, 36). However, the structure of the free RNA may not be the same as that bound to Lin28 because our data suggest that Lin28 unwinds G4 quartets. This is consistent with footprinting assays of pre-let-7g, which previously suggested that Lin28 binding induces a conformational change within the RNA (32). Isolated studies with the ZnF domains also suggest that they partially unfold pre-let-7 targets (56). Additionally, a FRET study revealed that the CSD of Lin28 remodels the terminal loop of pre-let-7 by unwinding the upper stem region (37). We propose that the Lin28-mediated remodeling of RNA disrupts the G4 structure because in the presence of Lin28, the Lin28-binding RNAs no longer induce NMM fluorescence.

A recent study found that another CSD protein, YB-1, binds to G4s in tRNA fragments (65). Truncation analysis showed that the CSD is responsible for G4 binding of YB-1. That study and our data suggest that the CSD, a nucleic acid-binding domain, which first arose in bacteria and forms an antiparallel β barrel, functions by binding and remodeling G4s. Further studies are needed to explore this hypothesis.

Remodeling G4s could be essential for Lin28 function (Fig. 6). During let-7 biogenesis, Lin28 recruits TUT4 to pre-let-7 transcripts and then TUT4 uridylates pre-let-7s to promote their degradation (28, 29). TUT4 does not bind to pre-let-7 in the absence of Lin28 (29). Remodeling of the pre-let-7 G4 by Lin28 could be required for the TUT4 interaction and subsequent uridylation. Because the G4 structure might interfere with ribosome scanning, remodeling G4s in mRNAs could also explain the ability of Lin28 to stimulate mRNA translation.

G-quadruplexes are found within DNAs in eukaryotic telomeres (66); the promoter regions of some cancer-related genes, including BCL2 (67), KIT (68), MYB (69), MYC (70), HIF1A (71), hTERT (72), KRAS (73), RB (74), and VEGF (75); immunoglobulin gene switch regions (76); and ribosomal genes (77). G4-intercalating agents have been assessed for anticancer ther-

![Figure 6. Model of Lin28 RNA regulation.](http://www.jbc.org/)

**Figure 6. Model of Lin28 RNA regulation.** A and B, we propose that Lin28 binds and remodels G4s in pre-let-7 miRNAs (A) and in LREs of mRNAs (B). Remodeling pre-let-7s might be required for TUT4 binding, subsequent uridylation, and degradation of pre-let-7s. Remodeling G4s in LREs may increase ribosome scanning and stimulate mRNA translation. The Lin28 CSD is in blue, and the ZnFs are in green.
apy (57, 50, 51). TMPyP4, a porphyrin analogue of NMM that binds G4s and double-stranded DNA, blocks MYC transcription and decreases tumor growth and increases survival in xenograft studies (57, 58). By blocking transcription and translation of oncogenes, G4-intercalating agents have demonstrated anticancer activity (57, 50, 51). Similar agents could be designed to target the G4 feature of Lin28-binding RNAs, preventing Lin28 remodeling G4s and inhibiting Lin28 activity.

Here, we have shown that NMM inhibits Lin28 binding to RNAs in a gel shift. This is the first example of a small-molecule inhibitor of Lin28. NMM may also inhibit Lin28 in cells. The addition of NMM to embryonal carcinoma cells increased let-7 miRNAs levels and decreased protein levels of Lin28 mRNA targets OCT4, HMGA1, CCNB1, CDK4, and Lin28. NMM also decreased the number of sphere-forming cells and inhibited colony formation. These findings suggest that NMM inhibits self-renewal. Because many human cancers, especially those with the worst prognosis, are often undifferentiated and have increased Lin28, NMM or other G4-intercalating agents may possess anticancer activity in these tumors. Our findings add new insight to understanding how Lin28 selects and regulates its miRNA and mRNA targets and suggest a new strategy for therapeutic intervention.

Author Contributions—E. O. conceived the study, conducted the experiments in Table 1, Fig. 2, Fig. 3, Fig. 4, Fig. 5, A, B, E, F, G, and H, and Fig. 6, and wrote the paper. M. T. N. L. performed experiments in Fig. 5, C and D. S. I. synthesized all the RNA constructs. S. M. T. performed experiments in Fig. 1C. R. K. and O. H. provided technical advice. H. A. helped to design and to execute the NMR and NMM experiments. G. W. and J. L. analyzed the data, coordinated the studies, and edited the paper. All authors reviewed the results and approved the final version of the manuscript.

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Elizabeth O'Day, Minh T. N. Le, Shunsuke Imai, Shen Mynn Tan, Rory Kirchner, Haribabu Arthanari, Oliver Hofmann, Gerhard Wagner and Judy Lieberman

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