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A Conditional System to Specifically Link Disruption of Protein-Coding Function with Reporter Expression in Mice

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SUMMARY

Conditional gene deletion in mice has contributed immensely to our understanding of many biological and biomedical processes. Despite an increasing awareness of nonprotein-coding functional elements within protein-coding transcripts, current gene-targeting approaches typically involve simultaneous ablation of noncoding elements within targeted protein-coding genes. The potential for protein-coding genes to have additional noncoding functions necessitates the development of novel genetic tools capable of precisely interrogating individual functional elements. We present a strategy that couples Cre/loxP-mediated conditional gene disruption with faithful GFP reporter expression in mice in which Cre-mediated stable inversion of a splice acceptor-GFP-splice donor cassette concurrently disrupts protein production and creates a GFP fusion product. Importantly, cassette inversion maintains physiologic transcript structure, thereby ensuring proper microRNA-mediated regulation of the GFP reporter, as well as maintaining expression of nonprotein-coding elements. To test this potentially generalizable strategy, we generated and analyzed mice with this conditional knockin reporter targeted to the Hmga2 locus.

INTRODUCTION

In recent years, methods allowing manipulation of genes of interest within living organisms have enabled a detailed understanding of numerous genes’ function across diverse phyla. The ability to alter the mouse genome has been especially critical for the investigation of gene function in vivo and has contributed immensely to our understanding of many fundamental questions in developmental biology and biomedical sciences. In particular, the development of site-specific recombinase systems in mice, including the Cre/loxP and FLP/FRT systems, has allowed the inactivation of genes of interest with both spatial and temporal control. As a result, conditional gene disruption has become a critical tool for understanding gene function during development, homeostasis, as well as in disease states (Rajewsky, 2007; Schmidt-Supprian and Rajewsky, 2007). Fluorescent reporter alleles have proved to be another key resource in the dissection of gene function by allowing direct visualization and isolation of molecularly defined subsets of cells (Hadjantonakis et al., 2003).

The past decade has brought an increased awareness that protein-coding genes can also possess additional nonprotein-coding functions. Single transcriptional units with multiple functions suggest interesting mechanisms that influence the cellular state, but their very existence necessitates new genetic methods to specifically abrogate the protein-coding element without altering other functions of the transcript and to generate reporter alleles that remain subject to all endogenous forms of regulation. In developing an allele system to couple specific disruption of protein-coding function with reporter expression, we considered the potential noncoding functions of transcripts.

Approximately half of all microRNAs (miRNAs) are encoded within introns. Targeted deletion of several intronic miRNAs has resulted in severe phenotypes including embryonic lethality (Kuhnert et al., 2008; Miyaki et al., 2010; Nakamura et al., 2011; Wang et al., 2008; Zhao et al., 2007), whereas many other intronic miRNAs remain uncharacterized. Therefore, gene targeting that inadvertently disrupts the expression of intronic miRNAs could have unanticipated phenotypic consequences (Kuhnert et al., 2008; Osokine et al., 2008; Wang et al., 2008). Interestingly, rare exonic miRNAs have also been shown to have critical functions (Sundaram et al., 2013).
Further indicating the need for improved gene inactivation methods, recent work indicates that miRNAs may also directly influence miRNA function through competitive binding, suggesting that the relative abundance of one miRNA target may impact that miRNA’s ability to repress other targets. Genetic deletion or truncation of transcripts that have this competitive endogenous RNA (ceRNA) activity could thus lead to inadvertent widespread alterations in the protein expression of conserved miRNA targets (Karreth et al., 2011; Kumar et al., 2014; Poliseno et al., 2010; Salmena et al., 2011; Tay et al., 2011). More generally, any titratable site-specific RNA-binding factor could be affected by the deletion of a highly expressed target transcript. Finally, recent reports of abundant circular RNA species comprised of exons of protein-coding genes with potential additional activity forewarn of further inadvertent changes in cellular state when transcripts are entirely ablated or truncated (Memczak et al., 2013; Salzman et al., 2012).

Additionally, the appropriate regulation of protein expression from reporter alleles can also be influenced by transcript structure. miRNAs, RNA-binding proteins, and RNA secondary structure regulate transcript stability and protein expression. Thus, reporters that are embedded within the native transcript will more faithfully recapitulate the expression of the gene of interest. Indeed, reporter alleles with endogenous versus exogenous 3’ UTRs can produce dramatically different expression patterns due to the presence or absence of appropriate miRNA-mediated regulation (Merritt et al., 2008; Yoo et al., 2009).

Collectively, the potential for transcripts to possess multiple functions indicates that more precise genome modification methods are required to avoid the potential phenotypic consequences of altering nonprotein-coding functions of genes. Based on these considerations, we developed a murine allele system in which conditional disruption of protein-coding function is directly coupled to fluorescent marker expression without impacting other aspects of transcript function.

**RESULTS**

**A Strategy to Couple Conditional Gene Disruption with GFP Reporter Expression in Mice**

The alteration of gene function with temporal and spatial control as well as the use of fluorescent reporter alleles has allowed the detailed investigation of a plethora of developmental processes. Coupling conditional Cre/loxP-mediated gene disruption with fluorescent reporter expression would not only generate two useful tools from one gene-targeting event but would also allow single-cell resolution of gene disruption, permit altered cells to be assessed relative to nontargeted cells, and enable the identification and isolation of cells that are attempting to express the disrupted gene.

Several genetic approaches in mice have been previously developed to inactivate a gene of interest and express a reporter gene, including the combination of Cre reporter alleles with floxed alleles (Alexander et al., 2009; Lao et al., 2012; Srinivas et al., 2001), mosaic analysis with double markers (Tasic et al., 2012; Zong et al., 2005), systems based on the inversion of gene-trap-like elements (Mandalos et al., 2012; Schnütgen et al., 2003, 2005; Schnütgen and Ghyselinck, 2007; Xin et al., 2005), and floxing of the entire gene followed by a 3’ reporter (Moon et al., 2000; Potocnik et al., 2000; Theis et al., 2001; Wellerhaus et al., 2008). Each of these methods has its unique strengths, but system-specific limitations have in all cases negated their widespread application (Table S1).

We devised a strategy based on the insertion of an inverted splice acceptor-eGFP-splice donor (SA-GFP-SD) cassette into an intron of the target gene. Pairs of heterologous loxP sites (2272 loxP and 5171 loxP) (Lee and Saito, 1998; Stern et al., 2008) flank the SA-GFP-SD cassette, thus allowing two sequential Cre-mediated recombination events to stably invert the SA-GFP-SD cassette. The inversion should disrupt gene function and concomitantly generate a GFP reporter (Figures 1A and S1). The first recombination event inverts the intervening sequence between either the 5171 loxP or the 2272 loxP sites and generates one of two intermediates in which alternate loxP variants are now in a head-to-tail orientation. The second recombination event then deletes the sequence between these sites leaving a final stably inverted cassette with one 2272 loxP site and one incompatible 5171 loxP site (Figures 1A and S1). Most importantly, we engineered a splice donor site following the stop codon of GFP to allow the transcript to continue through the remaining exons and 3’ UTR to the endogenous polyadenylation sites (Figures 1A and 1B). Because the reading frame at the end of the first exon of a gene of interest can be in any of the three reading frames, we generated base vectors with GFP in each reading frame to generate this style of conditional knockin (CK) allele (Figure S1).

**Generation of an Hmga2CK Allele**

To test our strategy to link Cre-mediated gene disruption with GFP reporter expression, we generated a CK allele of the chromatin-associated protein Hmga2 (Hmga2CK). We chose this gene because of our interest in Hmga2 in cancer metastasis as well as its dramatic role in body size regulation and potential role in stem cell biology (Nishino et al., 2008; Winslow et al., 2011; Xiang et al., 1990; Yu et al., 2007; Zhou et al., 1995). The Hmga2 genomic locus is quite large (~116 kb), has all exons in the same frame, has an intronic miRNA (miR-763), and produces a transcript that is highly regulated by endogenous miRNAs. Despite the difficulty of accounting for these common elements with existing genetic-targeting approaches, these characteristics made Hmga2 an excellent candidate for our CK allele system. The Hmga2 locus was targeted with the SA-GFP-SD cassette in the reverse orientation (Figure 1A). Mice generated from correctly targeted embryonic stem (ES) cells were bred with mice expressing FLPe recombinase to delete the neomycin-resistance cassette, thereby generating the Hmga2CK allele (Farley et al., 2000) (Figure 1A). To characterize the Hmga2 allele after cassette inversion, Hmga2CK mice were then bred with mice expressing Cre recombinase to generate a germline Hmga2GFP allele (Wellerhaus et al., 2008) (Figure 1A).

**Cre-Mediated Stable Inversion of the SA-GFP-SD Cassette Disrupts Protein Expression**

The Hmga2GFP allele generated a transcript in which exon 2 was replaced by GFP, resulting in a protein product containing the N terminus of Hmga2 fused to GFP (Figures 1B–1D and S1). Although the SA within our cassette is well-characterized
and is capable of generating null-like alleles (Carette et al., 2009), it remained possible that some Hmga2 transcripts from the Hmga2GFP allele could splice from exon 1 to exon 3. In Hmga2GFP/GFP embryos and murine embryonic fibroblasts (MEFs), no evidence of an mRNA generated from exon 1 splicing to exon 3 was detected by RT-PCR (data not shown). Additionally, whereas splicing from exon 1 to exon 3 would create an in-frame protein product, even very long exposure times gave no indication of any truncated Hmga2 protein product in Hmga2GFP/GFP MEFs, or western blot analysis of embryos and MEFs (Figures 1D, S2A, S2B, and S3C; data not shown). To verify the conditional nature of the allele, we infected Hmga2CK/+ embryos with an adenoviral vector expressing Cre. Cre expression converted the CK alleles to their GFP conformation, with infected cells expressing high levels of GFP and lacking full-length Hmga2 protein (Figures 2F and S3C).

Cassette Inversion Maintains the Physiologic Expression of Intronic miRNAs
Approximately half of known miRNAs are encoded within the introns of protein-coding genes. Although some gene inactivation strategies disrupt transcript expression, thereby disrupting both the protein-coding mRNA and the intronic miRNA, our system maintains the production of the full-length transcript, suggesting that intronic miRNAs should not be affected (Figures 1A and 1B). miR-763 resides within intron 3 of the Hmga2 locus; therefore, we quantified miR-763 expression in MEFs and embryos carrying the Hmga2CK and Hmga2GFP alleles. Importantly, neither the Hmga2CK nor the Hmga2GFP allele affected miR-763 expression, confirming that our gene-targeting strategy can specifically inactivate the protein-coding element without impacting other functional components within downstream introns (Figures S2I–S2L).

Cassette Inversion Maintains the Full-Length Transcript Allowing Physiologic miRNA-Mediated Regulation of the GFP Reporter
One theoretical advantage of generating a conditional allele using our general strategy is the ability to embed the GFP reporter
within an otherwise full-length transcript, thereby ensuring that the GFP reporter remains under all the transcriptional and post-transcriptional regulatory control elements of the endogenous locus. In particular, the inclusion of a splice donor site after the stop codon of GFP allows transcription to continue all the way through the 3' UTR, suggesting that the conditional GFP reporter should remain under physiologic miRNA regulation (Figures 1A, 1B, S1H, S3B, and S3D). To directly assess whether the Hmga2GFP allele maintains a physiologic and full-length transcript structure, we performed northern blot analyses using Hmga2- and GFP-specific probes. The composite Hmga2GFP transcript was detected as a single predominant band that was similar in size to the Hmga2wt counterpart, indicating that cassette inversion embeds GFP within the full-length transcript (Figures S3E and S3F).

Because Hmga2 is a well-characterized let-7 miRNA target (Lee and Dutta, 2007; Mayr et al., 2007) and is regulated by let-7 in MEFs (Viswanathan et al., 2008; Figures 3C and S4), we
Figure 3. Hmga2Ex1-GFP Remains under miRNA Control

(A) Flow cytometric analysis of Hmga2GFP/GFP MEFs after transfection with a let-7 mimic documents let-7-mediated reduction in GFP fluorescence. (B) Quantification of the GFP mean fluorescence intensity (MFI) after transfection with a control miRNA mimic or increasing amounts of let-7 mimic (2, 10, and 36 pmol; only 36 pmol let-7 mimic is shown for Hmga2<sup>+/+</sup> cells). Mean ± SD of triplicate wells is shown. Data are representative of four experiments. (C) Let-7-mediated Hmga2Ex1-GFP protein reduction parallels that of full-length Hmga2 in Hmga2<sup>GFP/GFP</sup> MEFs. Cells were transfected with a control miRNA mimic (36 pmol) or increasing amounts of let-7 mimic (2, 10, and 36 pmol). Both Hmga2 and Hmga2<sup>Ex1</sup>-GFP expression decreases in a dose-dependent manner. A nonspecific band (n.s.) is marked. Actin shows equal loading. short exp., short exposure; long exp., long exposure. (D) Quantification of GFP MFI after stable retroviral Lin28 expression. Mean ± SD of triplicate wells is shown.

performed a series of experiments to test whether the Hmga2<sup>GFP</sup> allele remains under let-7 control. Transfection of Hmga2<sup>GFP/+/+</sup> or Hmga2<sup>GFP/+/+</sup> MEFs with a synthetic let-7 mimic reduced GFP expression in a dose-dependent manner (Figures 3A–3C, S4A, and S4B; data not shown). Furthermore, the reduction of Hmga2Ex1-GFP protein mirrored that of endogenous Hmga2 after transfection of Hmga2<sup>GFP/+</sup> MEFs with the let-7 mimic (Figure 3C). The Hmga2–GFP reporter was also increased by stable expression of the let-7 inhibitor Lin28 (Figures 3D and S4C). Collectively, these results indicate that the Hmga2Ex1-GFP reporter generated after Cre-mediated cassette inversion remains under appropriate posttranscriptional 3’ UTR regulation.

Cassette inversion introduced a premature stop codon after the GFP-coding sequence followed by several additional exons; therefore, it is possible that the composite transcript could be subject to nonsense-mediated decay (NMD) (Chang et al., 2007). To directly compare the expression of the Hmga2<sup>Ex1</sup> and Hmga2<sup>GFP</sup> transcripts in isogenic cells, we treated Hmga2<sup>CK/CK</sup> cells with a Cre-expressing adenoviral vector to generate Hmga2CK<sup>GFP</sup> cells. Quantitative PCR and northern blotting for total Hmga2 indicate that the expression of Hmga2<sup>Ex1</sup> in Hmga2<sup>CK/GFP</sup> cells is comparable to the expression of Hmga2<sup>GFP</sup> in Hmga2<sup>GFP/GFP</sup> cells. Additionally, inhibition of NMD induced comparable upregulation of Hmga2<sup>GFP</sup> and Hmga2<sup>Ex1</sup> transcripts, had no effect on let-7-mediated repression of either Hmga2<sup>Ex1</sup> or Hmga2<sup>GFP</sup> mRNA or protein, and did not alter Lin28-induced upregulation of Hmga2<sup>GFP</sup> protein (Figure S4).

**Hmga2 Protein Is Responsible for the Phenotypes Associated with Hmga2 Deficiency**

Previously, Hmga2 null alleles have been generated through spontaneous deletion, fortuitous insertional mutagenesis, and conventional gene targeting. In all cases, Hmga2 null mice displayed a dramatic dwarfism phenotype (Xiang et al., 1990; Zhou et al., 1995). To determine whether Hmga2<sup>GFP</sup> mice phenocopy Hmga2 null mice, we analyzed the growth of Hmga2<sup>+/+</sup>, Hmga2<sup>GFP+</sup>, and Hmga2<sup>GFP/GFP</sup> mice. Consistent with the Hmga2 null phenotype, postnatal day 0 (P0) Hmga2<sup>GFP/GFP</sup> mice were slightly smaller than Hmga2<sup>+/+</sup> and Hmga2<sup>GFP/+</sup> littermates (Figure 4A). Both male and female Hmga2<sup>GFP/GFP</sup> mice recapitulated the expected dwarfism phenotype, with 12-week-old mice being shorter and ~50% the weight of control mice (Figures 4B–4F). Additional aspects of the Hmga2 null phenotype were also observed in the Hmga2<sup>GFP/GFP</sup> mice, including a disproportionate reduction in fat tissue and pinnae size and a relatively mild reduction in brain size (Nishino et al., 2008) (data not shown). Hmga2<sup>CK/GFP</sup> mice were phenotypically indistinguishable from Hmga2<sup>GFP/+</sup> control mice, indicating that the cassette itself does not dramatically affect the expression of the targeted allele prior to its inversion by Cre (Figures S2E and S2F).

**DISCUSSION**

The unequivocal coupling of gene disruption and reporter expression, unaltered expression of intrinsic miRNAs in the presence of disrupted protein-coding function, and the induction of a fluorescent reporter under the control of all endogenous genomic elements and 3’ UTR control regions all support the broad use of CK alleles to better understand gene function in vivo. Because miRNA-mediated control of protein expression and the role of ceRNA networks are almost completely unstudied in this context, combined approaches that employ both CK alleles and alleles that entirely abrogate transcript expression could represent an important future method to study the importance of ceRNA and other noncoding transcript functions in complex tissues and at the organismal level. This system also circumvents the major difficulties encountered with mosaic deletion resulting from ineffective Cre-ER or weak cell-type-specific Cre alleles because it allows cells that harbor the deleted target gene to be definitively distinguished from those that do not and, in fact, allows for neighboring control and null cells to be
analyzed within the same tissue. Although other strategies to link gene disruption with reporter expression have been developed, none allows the generalizable and streamlined generation of alleles that specifically disrupt protein-coding function while leaving all nonprotein-coding functions intact (Table S1). The CK allele system will be widely applicable for creating Cre-regulated alleles to better understand the protein-coding element of genes during many stages of development, normal homeostasis, and disease states.

The CK allele system should be amenable for targeting most genes because it employs a well-characterized SA that has been used for multiple genome-wide loss-of-function screens in haploid cells (Bürckstümmer et al., 2013; Carette et al., 2011). However, a few situations do exist that preclude the use of the CK technology. Introns flanked by noncanonical SA-donor pairs are rare, constituting less than 1% of introns, but would be insensitive to an inverted CK allele. Additionally, our system is also limited to genes that contain at least one intron and in which an exon 1-GFP fusion would lack function.

mRNA stability is influenced by RNA sequence and structure, and both are altered in the CK allele after cassette inversion and incorporation of the GFP sequence. Additionally, interrupting the protein-coding function by introducing a premature stop codon after GFP could induce NMD (Brogna and Wen, 2009).

Although we found that switching of the Hmga2wt transcript to Hmga2GFP did not reduce steady-state mRNA abundance, the effect of cassette inversion on transcript stability on additional targets will have to be determined empirically.

Hmga2 is an important regulator of body size across a wide range of species, including humans (Makvandi-Nejad et al., 2012; Weedon et al., 2007). Previous murine Hmga2 null alleles have all resulted from the abrogation of the entire Hmga2 transcript (Xiang et al., 1990; Zhou et al., 1995). That our Hmga2GFP allele recapitulates the null phenotypes confirms that most, if not all, of the developmental phenotypic consequences observed in these earlier models are driven by the loss of Hmga2 protein function rather than the function of miR-763 or disruption of a let-7-Hmga2 ceRNA network. Our results provide a loss-of-function complement to the increased body size in transgenic mice expressing increased levels of Hmga2 protein (Arlotta et al., 2000; Battista et al., 1999). Given that the molecular functions of Hmga2 during development and disease remain poorly characterized, our allele should aid in dissecting the role of this important chromatin-regulating protein.

EXPERIMENTAL PROCEDURES

Creation of Base-Targeting Vectors

The allele design (termed the CK system) is based on the creation of an in-frame GFP fusion protein; therefore, CK plasmids were generated for each of the three reading frames (Figure S1; pCK [+0], pCK [+1], and pCK [+2]). To generate these plasmids, eGFP-SA-2272loxP-5171loxP fragments in each of the three reading frames were amplified with the addition of a 5' multiple cloning site, a consensus splice donor site (in reverse, actcacctt), and a 3' PacI site. These fragments were ligated into a plasmid backbone that contained a diphtheria toxin A expression cassette, a consensus splice donor site (in reverse, actcacctt), and a 3' PacI site. These fragments were ligated into a plasmid backbone that contained a diphtheria toxin A expression cassette, a multiple cloning site, an FRT-flanked neomycin-resistance gene (neo), 2272loxP-5171loxP sites, and a final multiple cloning site (Figures 1A and S1; Supplemental Experimental Procedures).
Generation of CK Alleles

Construction of a CK allele for any gene of interest requires the amplification and ligation of the targeting arms into the appropriate pCK plasmid. Having an exon within the region of the genome to be inverted is not required for the system to work, but an exon can be included if desired (as in the Hmga2CK allele). Reasons to incorporate an exon may include the ATG start site lies within the second exon, a potential alternate ATG start lies within the second exon, or if one simply desires to further guarantee gene disruption. If no exon is included, it would likely be preferable to insert the cassette in the first intron of the target gene to limit the length of the endogenous gene produced prior to splicing into the SA-GFP-splice donor cassette after Cre-mediated inversion. In cases where an exon is included within the region that will be inverted, the overall conformation would likely be very similar to the Hmga2CK allele that we have generated. If an exon is to be included, it will be necessary to clone three regions into the pCK vector in any order; otherwise only two regions will be required (Figure S1A; Supplemental Experimental Procedures).

Hmga2 Targeting and Generation of Hmga2CM and Hmga2zpr Mice

The Hmga2-targeting construct was linearized and electroporated into ES cells (a kind gift from Rudolph Jaenisch) using standard conditions. Neomycin (300 μg/ml G418)-resistant colonies were picked, expanded, and screened by long-range PCR for correct targeting of the Hmga2 locus. Of 294 clones, 5 (1.7%) were correctly targeted. C57BL/6J blastocystics were injected with targeted ES cells resulting in several high-percentage chimeras. Targeted Hmga2zpr mice were bred to Rosa26LP/Lp mice (The Jackson Laboratory; stock number 003946) to delete the neo cassette and generate the Hmga2CK allele. A subset of Hmga2zpr mice was bred to β-actin-Cre mice to create mice with a germline Hmga2zpr allele (Figure 1A). The MIT and Stanford Institutional Animal Care and Use Committees approved all animal studies and procedures.

Generation of MEFs, Adenoviral Infection, and Flow Cytometry

MEFs were generated from embryos between E12.5 and E16.5. To express Cre recombinase in MEFs, cells were infected with either Adeno-Cre or Adeno-FLPo (control) purchased from the Gene Transfer Vector Core at the University of Iowa. Analysis of the cells was performed 3 days after infection. Data were collected on a BD LSR II analyzer (BD Biosciences) at the Stanford Shared FACS Facility. For the quantification of GFP in MEFs on a LSR II flow cytometer, cells were thoroughly trypsinized and stained with DAPI to exclude dead cells.

Western Blot Analysis

MEFs were trypsinized and pelleted before being lysed. Denatured samples were separated by SDS-PAGE, followed by transfer to polyvinylidene fluoride membranes. Membranes were stained with the indicated primary antibodies rabbit anti-Hmga2-P1 (BioCheck; 59170AP), rabbit anti-GFP (Cell Signaling Technology; 2936), and β-actin antibody (Sigma-Aldrich; A5441). For Li-COR imaging, membranes were first stained with the indicated primary antibodies followed by staining with either the goat anti-mouse-IRDye 800CW (926-32210; Li-COR Biosciences) or the goat anti-rabbit-Alexa Fluor 680 (A-21109; Life Technologies) for 1 hr at room temperature in the dark. After three washes (1× PBS, 0.05% Tween 20), the membranes were imaged with the Li-COR imager Li-COR Odyssey.

Upf1 Knockdown

The small interfering RNA (siRNA) specific for murine Upf1 transcripts was purchased from Life Technologies (siRNA ID, s72879). The procedure to deliver the control and Upf1 siRNA into MEFs is described below. Transfectants were harvested 3 days posttransfection and lysed for quantification of mRNA. For functional validation of Upf1 knockdown, Gas5 transcript was used as described by Keeling et al. (2013).

Northern Blot Analysis

Northern blots were performed using standard methods. In brief, radiolabeled probes were generated through random hexamer-primed synthesis by using a PCR-amplified cDNA sequence as the template and DNA polymerase I Klenow fragment and α-32P-dCTP. Probes were purified using a spin column kit (Nucleotide Removal Kit; QIAGEN). Membranes were prewashed, hybridized with radiolabeled probe for 16–20 hr at 42°C, washed twice with 2× saline sodium citrate (SSC) and 0.1% SDS at 42°C for 20 min, and twice with 0.1× SSC and 0.1% SDS at 55°C for 20 min. Membranes were imaged on a Molecular Dynamics Typhoon 9400 Imager (Amersham/GE Healthcare).

Let-7 Transfection and Retroviral Lin28 Expression

To deliver control (mirVana-negative control mimic) and/or let-7 (mirVana mmu-let-7a-5p miRNA mimic) miRNA mimics into MEFs, Lipofectamine RNAiMAX was used according to the manufacturer’s description (Life Technologies). To normalize total input mimics, compound levels of control mimic were added with titrated let-7 mimic. Overexpression of the let-7 inhibitor Lin28 (Viswanathan et al., 2009) was achieved by transducing MEFs with a retroviral vector (murine stem cell virus [MSCV]) harboring the Lin28 cDNA (Addgene; pMSCV-mLin28a/neo, plasmid No. 26357) (Viswanathan et al., 2009). An MSCV-neo vector without Lin28 was used as a negative control. Transduced MEFs were cultured in the presence of G418 (0.5 mg/ml) for 3 days to allow complete enrichment of transductants. Selected MEFs were assessed for GFP and Hmga2 levels using flow cytometry and western blot analysis.

SUPPLEMENTAL INFORMATION

Supplemental Information includes Supplemental Experimental Procedures, four figures, and one table and can be found with this article online at http://dx.doi.org/10.1016/j.celrep.2014.05.031.

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