MiR-146a wild-type 3’ sequence identity is dispensable for proper innate immune function in vivo

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The prevailing model of microRNA function is that the “seed region” (nt 2–8) is sufficient to mediate target recognition and repression. However, numerous recent studies have challenged this model, either by demonstrating extensive 3’ pairing between physically defined miRNA–mRNA pairs or by showing in Caenorhabditis elegans that disrupted 3’ pairing can result in impaired function in vivo. To test the importance of miRNA 3’ pairing in a mammalian system in vivo, we engineered a mutant murine mir-146a allele in which the 5’ half of the mature microRNA retains its wild-type sequence, but the 3’ half’s sequence has been altered to robustly disrupt predicted pairing to this latter region. Mice homozygous or hemizygous for this mutant allele are phenotypically indistinguishable from wild-type controls and do not recapitulate any of the immunopathology previously described for mir-146a–null mice. Our results indicate that 3’ pairing is dispensable for the established myeloid function of this key mammalian microRNA.

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Introduction

miRNAs are well-established regulators of mRNA stability and translation (Bartel, 2018). The discovery that hundreds of distinct miRNAs are expressed in animals (Lagos-Quintana et al, 2001; Lau et al, 2001; Lee & Ambros, 2001) and the subsequent finding that, unlike plant miRNAs, bilaterian animal miRNAs were only rarely predicted to base pair to their regulatory targets with full complementarity (Mansfield et al, 2004) raised the challenge of predictively identifying miRNA targets via computational analysis. Seminal work by the Burge and Bartel laboratories revealed that the most highly conserved pairing between miRNAs and mRNA 3’ UTRs was predicted to occur between nt 2 and 8 of mammalian miRNAs and their mRNA targets (Lewis et al, 2003). This finding was broadly incorporated into a generation of miRNA target prediction algorithms, and several subsequent studies corroborated these initial predictions by showing that the interaction between previously established miRNA–mRNA pairs was not only dependent upon the seed sequence but, moreover, independent of or only marginally reliant on non-seed sequence (Doench & Sharp, 2004; Kloosterman et al, 2004; Brennecke et al, 2005; Lai et al, 2005; Lim et al, 2005). The first unbiased identification of bona fide miRNA/mRNA target pairs by physical association of Argonaute proteins with mRNA (Chi et al, 2009) corroborated the predictions of these existing algorithms to some extent by showing that most identified targets contained a seed match, a result which would be echoed by subsequent studies using similar methodologies (Hafner et al, 2010; Chi et al, 2012; Loeb et al, 2012; Grosswendt et al, 2014; Moore et al, 2015; Broughton et al, 2016). Some of these latter studies would also establish that target sites that contained a seed were typically more effective than sites that did not (Chi et al, 2012; Loeb et al, 2012; Moore et al, 2015). In addition to this, however, these studies also showed that the vast majority of these Argonaute/mRNA interactions also incorporated binding beyond the seed, within the 3’ region, and a significant proportion of microRNA response elements (MREs) contained no identifiable seed match at all. This growing body of literature has lent support to the notion that various degrees of base pairing between miRNA 3’ sequences and associated mRNA targets is the rule, rather than the exception. A functional role for 3’ pairing has also recently been corroborated in vitro in mammalian cells (Jeong et al, 2017) and in vivo in Caenorhabditis elegans (Zhang et al, 2015; Broughton et al, 2016; Brancati & Großhans, 2018). The collective implication of these studies is that the 3’ region of a given miRNA may confer an important but not-yet-fully defined role in determining the target spectrum (and thus function) of the miRNA. However, this has not previously been genetically addressed in vivo in a mammalian system.

We thus constructed a “flipped” allele of an miRNA whose loss-of-function has been previously established to yield a robust
phenotype in mice. Within this allele, the sequence of the 3′ half of the mature miRNA was exchanged with that of its complementary strand sequence in the pre-miRNA hairpin. We chose for our model, miR-146a, a pivotal immunoregulatory miRNA. MiR-146a is one of the two members of the mir-146 family, the other being mir-146b, which differs from miR-146a in its mature sequence by only two nucleotides in the 3′ region. Despite this similarity, the two miRNAs are not functionally redundant in vivo; mice deficient for mir-146a but retaining wild-type mir-146b function are hypersensitive to LPS challenge and greatly predisposed to the development of hyperimmunity and myelodysplasia (Boldin et al, 2011). Given this robust phenotype, we bred mice homozygous for the “3′ flip (3′F)” allele to test how disruption of the 3′ region of this particular miRNA might impact miR-146a’s established genetic function. Closely following the workflow initially described for the characterization of mir-146a–deficient mice (and using the line as a control), we directly compared the phenotype of the mir-146a3′F, mir-146a-null, and wild-type alleles, both in the context of homozygosity and heterozygosity.

Results

MiR-146a 3′ pairing contributes to the regulation of target UTRs in vitro

We first used in vitro reporter assays to determine whether, in this context, miRNA “sensors” responded to ectopic miR-146a expression in a manner similar to UTRs that had previously been established to be miR-146a targets. To this end, we designed two synthetic miRNA duplexes: one identical to the duplex produced after cleavage of the wild-type mmu-mir-146a pre-miRNA by Dicer and the other designed after the theoretical 3′F duplex (Fig S1A). The mutant sequence was modeled to reflect the mature strand of a pre-miRNA in which each nucleotide from positions 13 to 20 of the 5′ strand was exchanged with the opposite nucleotide from the 3′ strand and vice versa. We reasoned that this mutation would be the most deleterious to any potential physiological pairing by this region of the mature miRNA by making the 3′ region of the mature miRNA as near to its complement as possible without disrupting the secondary structure of the hairpin. Positions 9–12 within the mature miR-146a were left unperturbed under the rationale that these sequences were likely to be unpaired in a physiological setting (Schirle et al, 2014; Bartel, 2018), and positions 21–22 were conserved to lessen the chance that changes to loop-adjacent sequences might impact the processing or regulation of the resulting mutant.

We constructed luciferase reporter constructs containing six tandem MREs specifically tailored to match either of these mimics within the pRL-TK-CX6X backbone (Doench et al, 2003). Thus, both the WT and 3′F mimics would perfectly base pair to their cognate reporter (with the exception of nt 10–13 to eliminate “slicer” activity), but only base pair to their non-cognate reporter via nt 1–9 (Fig S1B). We transfected HeLa cells with these reporters, challenging each reporter separately with a series of concentrations of both the wild-type and mutant siRNA mimic duplexes. In both cases, transfection of the mimic designed to pair to the reporter via both its seed region and the 3′ region repressed reporter activity to a significantly greater degree than the mimic designed to pair only via the seed region (Fig S1C and D, P-value < 0.0001, two-way ANOVA). These data demonstrated that extensive 3′ pairing has the potential to confer additional efficacy to miRNA function. However, because conventionally designed miRNA reporters, such as those assessed here, permit (or preclude) 3′ pairing to a more extreme extent than is thought to generally occur in a physiological setting, we next asked whether a similar phenomenon might be observed using reporters containing the 3′ UTRs of established miR-146a targets.

We thus repeated the reporter assay, this time targeting the 3′ UTRs of the two originally characterized targets of miR-146a, Irak1, and Traf6 (Taganov et al, 2006). Alignment of the 3′F miR-146a sequence with the known MREs in both targets predicted substantial disruption of 3′ base pairing (Fig S2A and B). In keeping with this, we observed a modest but statistically significant (P < 0.0001, two-way ANOVA) reduction in the ability of the 3′F mimic to repress both the Irak1 and Traf6 3′ UTRs (Fig S2A and S3A), whereas neither allele was capable of repressing mutated versions of the 3′ UTRs in which the central four seed nucleotides of each MRE had been mutated to their complement (Fig S2E and S2F). Taken together, these results support the notion that the 3′ non-seed region of miR-146a positively contributes to target repression in vitro.

Construction and validation of a murine mir-146a3′F allele

To determine whether the engineered “flip” mutation would be compatible with processing and expression from endogenous context, we synthesized and then subcloned the wild-type and 3′F miR-146a precursor hairpin sequences into the 3′ UTR of the pRL-TK-CX6X expression vector. Transfection of this vector into HEK 293 cells revealed processing and production of pre-miRNA and mature miRNA species from the mutant construct at absolute levels essentially identical to those observed for a similarly engineered wild-type miR-146a precursor construct in this context (Fig 1A and data not shown). We, therefore, targeted the endogenous mir-146a allele in murine embryonic stem cells via homologous recombination to replace the wild-type mmu-mir-146a locus with the mutant 3′F sequence (Fig 1B). In parallel, we deleted the mir-146a sequence in a second embryonic stem cell line, generating a null allele to serve as a positive control for mir-146a’s established phenotype (Fig 1C–F and data not shown). Mice derived from these embryonic stem cell lines and homozygous for either mutant allele were born at Mendelian frequency with no discernible overt phenotypes.

To confirm that the mir-146a3′F allele was expressed at similar levels to wild-type mir-146a, we compared the relative levels of mature wild-type and 3′F mir-146a in several tissues and immune lineages derived from homozygous animals using qRT-PCR. Absolute values for miRNA expression in each sample were calculated via comparison with serial dilutions of synthetic standards cognate to each predicted mature miRNA sequence, and cross-priming analysis revealed a minimum of ~200-fold specificity of each primer set for its respective target (Fig 1G). Absolute expression of the mature miR-146a3′F miRNA in mir-146a3′F/3′F homozygous mice
was similar to absolute expression of the wild-type mir-146a allele in homozygous wild-type mice in most tissues and cell populations examined (Fig 1H). As expected, neither wild-type nor 3’F mir-146a was detected in any tissues or cell populations derived from mir-146a−/− mice (Fig 1I and data not shown). Consistent with the qRT-PCR data, small RNA sequencing from BMDMs (Table S1) confirmed that mature miR-146a WT was processed at the correct register within cells derived from mir-146a WT/− mice (Fig S3A and B). Interestingly, these data also suggested higher levels of mature miR-146a in induced organs from mir-146a+/+ and mir-146a WT/− mice. Direct qRT-PCR quantitation (with standard) of WT mir-146a in indicated organs from mir-146a WT/− and mir-146a WT/− mice. Error bars = SD. In (H) and (I), CD4+ = CD4+ T cells; CD8+ = CD8+ T cells; BMDCs and BMDMds were either stimulated (+) for 24 h before lysis with LPS or else unstimulated (−).

Comparison of lifespan, hematology, and autoimmunity

We next set out to characterize mice homozygous for the mir-146a 3’F allele in comparison with mir-146a-null and wild-type animals, closely adhering to the workflow described in the initial published analysis of mir-146a deficiency (Boldin et al., 2011). As previously described, mir-146a−/− mice were marked by decreased long-term viability when compared with wild-type littermates, beginning at 6 to 8 mo of age (Fig 2A). Gross analysis of mir-146a−/− mice sacrificed during this period revealed marked splenomegaly (Fig 2B and C) and several signs of autoimmunity. Splenic T lymphocytes were characterized by increased basal activation on both CD4+ and CD8+ T lymphocytes, as defined by increased surface expression of CD69 and CD44 and decreased surface expression of CD62L. Spleens derived from mir-146a−/− mice were also characterized by an increase in GR-1+CD11b+ myeloid compartment size (Fig 2E and F), and the sera of these mice were marked by an elevated titer of anti-double-stranded DNA antibodies (Fig 2G). Also consistent with previous work, aged mir-146a−/− mice (~14 mo) were characterized by marked decreases in hematocrit, red blood cell count, and platelet (Fig 2H). Although we did not observe previously described reductions in the representation of white blood cells or hemoglobin at 14 mo of age (Boldin et al., 2011), our mir-146a−/− line exhibited significant increases in mean corpuscular volume, mean platelet volume, hemoglobin distribution width, and red cell distribution width (Fig S4), consistent with anemia and bone marrow failure. Histological analysis of the liver and kidney of aged mice from our mir-146a−/− line revealed increased infiltration of mononuclear inflammatory cells in both organs as compared with wild-type controls (Fig 2I and K). However, in contrast to what we had observed in vitro, none of these phenotypes, whether previously described or novel to this study, were observed in age-matched mir-146a WT/− mice. Indeed, mir-146a WT/− mice were phenotypically indistinguishable from wild-type controls (Fig 2A–K), whether...
at the previously described time points or when assessed again in the context of extremely advanced age (~125 wk, or roughly 2.2 yr, Fig 3).

To better rule out the possibility that the mir-146a 3'F allele was modestly hypomorphic, we repeated this battery of experiments, this time comparing mir-146a 3'F/hemizygotic mice with mir-146a +/+ controls. Once again, no statistically sufficient differences in phenotype were observed when these two lines were compared at the established time points, and no difference in longevity between the lines was observed out to 100 wk (Fig 4). Thus, for the phenotypes assessed, our data show that the mir-146a 3'F allele is functionally indistinguishable from the wild-type mir-146a allele.
MiR-146α-3′ sequence identity is dispensable for function

Our findings above did not rule out the possibility that some difference in function might be observed by comparing the miR-146α and miR-146α⁢3′ alleles in the context of challenge of the innate immune machinery, either in vivo or in vitro. We thus assessed this possibility, again closely following the previously established experimental workflow. As previously described, following sublethal challenge via injection of LPS (1 mg/kg) miR-146α-deficient mice exhibited elevated levels of serum TNFα, IL-1β, and IL-6 as compared with wild-type controls (Fig 5A). Analogous results were observed after challenge with a lethal (35 mg/kg) dose of LPS—miR-146α-deficient mice were characterized by elevated levels of serum TNFα, IL-1β, and IL-6 and succumbed to the lethal LPS dose at a significantly greater rate than wild-type controls (Fig 5B and C). However, although modest trends in cytokine production were observed in both LPS challenges, once again, responses in mice homozygous for the miR-146α⁢3′ allele were statistically indistinguishable from their respective controls at the level of cytokine production and viability (Fig 5A–C). Furthermore, repetition of these experiments with miR-146α⁢⁻/- and miR-146α⁢⁻/-/- mice revealed no significant differences in circulating cytokine levels or LPS sensitivity (Fig S5). Altogether, these results indicate that the miR-146α⁢3′ allele is essentially functionally equivalent to the wild-type miR-146α allele in these contexts.

To compare the function of the mutant and wild-type alleles in a cellular context, we challenged BMDMs from each line with LPS ex vivo. Again, consistent with previously published results, BMDMs from miR-146α-deficient mice were characterized by comparatively increased production of IL-6 at 8 and 16 h following the LPS challenge (Fig 5D). This response coincided with increased relative expression of IRAK1 protein beginning at 16 h post-challenge, consistent with the previously established role as a target of miR-146α (Fig 5E and F). Interestingly, we did not observe dysregulation of TRAF6 in any of our in vitro experiments (data not shown). Although we noticed a modest trend for increased production of IL-6 by LPS-challenged miR-146α⁢3′/-/- mice, this did not reach statistical significance (Fig 5D), and even this trend was significantly reduced when levels of IRAK1 protein were assessed (Fig 5E and F). Taken together, these data suggest that the miR-146α⁢3′ allele may be modestly hypomorphic compared with the wild-type allele but not to a degree that can be readily demonstrated.

Functional definition of miR-146α target spectrum

Having phenotypically assessed the function of the miR-146α allele within these contexts, we next wished to survey the transcriptome and proteome of miR-146α-deficient and miR-146α⁢3′/-/- BMDMs relative to wild-type controls. Three biological replicates of BMDMs derived from each of the three genotypes were stimulated with
1 ng/ml LPS for 16 h and harvested. Individual replicates were split into two fractions—one analyzed via next-generation sequencing and the other by label-free mass spectrometry proteome profiling.

Within mammalian systems, increased or decreased expression of a miRNA often correlates with a global shift in the relative expression of its predicted mRNA targets (Grimson et al., 2007). However, this characteristic shift was absent, not only in the mir-146a−/− BMDMs, but even, surprisingly, in the mir-146a−/− BMDMs when compared with wild-type controls. While differences of modest statistical significance were observed with lower target context scores (Grimson et al., 2007), the direction of these changes was opposite to what would be consistent with miRNA targeting, based on miRDB (Wong & Wang, 2014) was found to be up-regulated relative to wild-type BMDMs in the mir-146a−/− BMDMs, but even, surprisingly, in the mir-146a−/− BMDMs either (Fig 6B and Table S3). Indeed, only one protein—IRAK1—that was predicted by TargetScan 7.1 (Agarwal et al., 2015) and/or present within mirDB (Wong & Wang, 2014) was found to be up-regulated relative to wild-type BMDMs in the mir-146a−/− deficient cells. Of note, IRAK1’s expression was not significantly different between wild-type and mir-146a−/− F/B BMDMs (Fig 5E and F and Table S3).

Despite the fact that predicted miR-146a targets did not broadly increase in relative expression in mir-146a−/− BMDMs, we did identify several proteins that did, and also many whose expression decreased relative to wild-type controls (Table S3). We manually examined those proteins whose change in expression following LPS stimulation in the mir-146a−/− BMDMs differed from those in wild-type by a Z score greater than three, and we did not find any statistically significant enrichment for MREs complementary to miR-146a. This was true whether we looked for a perfect predicted target site at the proteomic level using mass spectrometry or segments of the coding sequence (as appropriate) of these genes of interest into the pRL-TK-CX6X reporter to test their responsiveness to miR-146a and miR-146a−/−. As a control, we also subcloned the respective 3’ UTRs or segments of the coding sequence (as appropriate) of these genes of interest into the pRL-TK-CX6X reporter to test their responsiveness to miR-146a and miR-146a−/− mimics. As a control, we also subcloned the respective 3’ UTRs of strongly up- and down-regulated genes which lacked any recognized miR-146a target site. None of the predicted targets or negative controls responded to either mimic (Fig 7A, multiple t tests, FDR = 1%). Similar results were obtained when we assessed 3’ UTRs of mRNAs containing miR-146a-complementary MREs and encoding proteins that differed in expression (Z score > 3) between wild-type and mir-146a−/− F/B BMDMs (Fig 7B). These
findings strongly argue against the possibility that differential protein expression of putative novel miR-146a and miR-146a-3’ targets is a direct effect of differential targeting by wild-type and mutant miRNA.

Discussion

We have characterized mutant mice in which the endogenous 3’ pairing of miR-146a, an established regulator of NF-κB and innate immune signaling, is disrupted. By carefully revisiting the battery of experiments initially used to characterize mir-146a–deficient mice, we demonstrate here that the mir-146a-3’ allele recapitulates the function of wild-type miR-146a in each of the previously established phenotypes. Some nonsignificant trends towards the mir-146a–deficient phenotype were noted in a comparison of mir-146a+/− and mir-146a-3’/− mice and cells, and we were able to document a modest difference in dose–response to reporters under the control of the Irak1 and Traf6 3’ UTRs. It is, thus, formally possible that the disruption of miR-146a’s endogenous 3’ pairing results in an allele that is very mildly hypomorphic, and that statistical significance would ultimately have been reached with greater numbers of mice. However, given that mir-146a-3’ mice thus far (100 wk at the time of writing) display indistinguishable health and longevity to mir-146a+/− littermates, any decreased function appears to be quite modest.

One might be tempted to consider that the mir-146a-3’ allele does, in fact, have impaired functionality, but that this is offset in mice of the relevant genotypes by increased relative expression/compensatory function of mir-146b, the sequence of which is highly similar to mir-146a. However, given that mir-146b wild-type function is genetically preserved in mir-146a–deficient mice, the phenotype associated with mir-146a deficiency is a product of the former’s wild-type function and the latter’s deficiency, respectively, and any genetic rescue of this phenotype can be most reasonably attributed to restoration of absent mir-146a function. Since the mir-146a-3’ allele rescues mir-146a deficiency to a level statistically indistinguishable from the wild-type allele (even within the hemizygotic state), we favor the conclusion that this allele, from a genetic standpoint and in the context examined, is essentially and functionally
equivalent to the wild-type mir-146a allele. Our small RNA sequencing data support this conclusion. Although miR-146b does appear to be elevated in cells derived from mir-146a–deficient mice, this elevation is not enough to maintain normal response to LPS stimulation in these cells, and no such elevation is observed in cells derived from mice homozygous for the mir-146a39F allele. If one were to speculate that this elevation of miR-146b in mir-146a–deficient mice was a product of homeostatic mechanisms pressuring compensatory function, one could then reasonably conclude that no such homeostatic pressure was present in the context of mir-146a39F allelic function.

Generally, our findings stand at apparent odds with both our initial in vitro data and with the large body of recent work demonstrating both that many miRNAs engage their targets via 3′ pairing and that disruption of this pairing can negatively impact the regulatory relationship between a given miRNA and its mRNA target (Chi et al., 2009; Hafner et al., 2010; Loeb et al., 2012; Helwak et al., 2013; Grosswendt et al., 2014; Moore et al., 2015; Broughton et al., 2016).

Regarding the former, we feel our work suggests that caution is warranted in the evaluation of in vitro models and, more specifically, that differences observable in the artificial context of reporter assays may not, in fact, accurately reflect or recapitulate in vivo function. Concerning the many studies where contribution of the miRNA 3′ region has been described, we do not feel that any of the experiments in our study directly challenge or invalidate the observations described in these previous works. Indeed, a consistent
the wild-type miR-146a sequences. Statistical comparisons were done using statistical significance as described in Fig S1 of these experiments, the control reporters to the far right are the same as those of the wild-type BMDMs. Left: proteins with reduced expression in response to LPS in the KO BMDMs; right: regions derived from proteins with reduced expression in comparison with the wild-type BMDMs. Left: proteins with reduced expression in response to LPS in the wild-type BMDMs as compared with the wild-type BMDMs right: regions derived from proteins with reduced expression in response to LPS in the KO BMDMs as compared with the wild-type BMDMs. Dual luciferase assays of putative miR-146a targets characterized by differential protein expression following LPS stimulation in a comparison of KO and wild-type BMDMs. Left: proteins with reduced expression in response to LPS in the wild-type BMDMs as compared with the wild-type BMDMs; right: regions derived from proteins with reduced expression in response to LPS in the KO BMDMs as compared with the wild-type BMDMs. In both of these experiments, the control reporters to the far right are the same as those described in Fig S1—synthetic reporters with six sequential MREs corresponding to the wild-type miR-146a sequences. Statistical comparisons were done using multiple t-test. For initial selection, false discovery rate was set at 1%. * denotes statistically significant differences unless noted. Error bars = SEM.

Figure 7. Targets differentially expressed during BMDM stimulation as a function of genotype are not direct miR-146a or miR-146a-3’ targets. (A) Dual luciferase assays of putative miR-146a targets characterized by differential protein expression following LPS stimulation in a comparison of KO and wild-type BMDMs. Left: proteins with reduced expression in response to LPS in the wild-type BMDMs as compared with the KO BMDMs; right: regions derived from proteins with reduced expression in response to LPS in the KO BMDMs as compared with the wild-type BMDMs. (B) Dual luciferase assays of putative miR-146a targets characterized by differential protein expression following LPS stimulation in a comparison of 3’ F and wild-type BMDMs. Left: proteins with reduced expression in response to LPS in the wild-type BMDMs as compared with the 3’ F BMDMs; right: regions derived from proteins with reduced expression in response to LPS in the 3’ F BMDMs as compared with the wild-type BMDMs. In both of these experiments, the control reporters to the far right are the same as those described in Fig S1—synthetic reporters with six sequential MREs corresponding to the wild-type miR-146a sequences. Statistical comparisons were done using multiple t-test. For initial selection, false discovery rate was set at 1%. * denotes statistically significant differences unless noted. Error bars = SEM.

Materials and Methods

Generation of mir-146a KO and 3’F mice

A ~7.1-kb region of the *Mus musculus* genome from chromosome 11 was amplified in two arms. Arm 1 (5.1 kb) was amplified from position 43,373,405 to 43,378,474 using primers 146_arm1_fw (5’-atcggagagctcccagg AACCGTGATCAGGGA6AAACG-3’) and 146_arm1_rev (5’-CACTTCAGCAGACCTGAAA-3’). Arm 2 (2.1 kb) was amplified from position 43,371,337–43,373,508 using primers 146a_arm2_FW (5’-atctgctgagctcAGAGACAAGAGATGGGCAATGTTGTTTG-3’) and 146a_arm2_rev (5’-atctgctgagctcAGAGACAAGAGATGGGCAATGTTGTTTG-3’). Arm 1, containing the mir-146a locus, was cloned into the HindIII site of the pIDT Smart ampicillin vector (IDT), and the WT mir-146a sequence was removed by cutting out a 536-nucleotide region flanked by a BspHI recognition site at the 5’ end and a Bcll recognition site at the 3’ end, then substituting in commercially synthesized sequences (IDT) either lacking the mir-146a pre-miRNA sequence entirely or comprising the 3’F mutation.

Arm 2 (5’ BamHI and 3’ NotI) and the edited Arm 1 (HindIII) were then subcloned into the pBluescript II KS(+) (Addgene) vector on either side of a loxP-flanked region containing the neomycin resistance cassette. The resulting mir-146a−3’F and mir-146a 3’ targeting...
constructs were electroporated into V6.5 murine embryonic stem cells, and the colonies were grown and picked under standard G418/ganciclovir double selection. Colonies derived from correctly recombined cells were identified via Southern analysis as described (Neilson et al., 2004), double digesting the genomic DNA with BamHI and EcoRV to detect insertion of an EcoRV site derived from the polylinker of the targeting vector. The Southern probes, amplified from genomic C57BL6.6 genomic DNA corresponded to chr11: 43,377,851–43,378,550 and chr11:43,370,951–43,371,356 (both GRCm38/mm10).

The G418 selection cassette was removed via transient transfection of Cre recombinase, and embryonic stem cells in which the cassette had been excised were identified via replicate plating in the presence and absence of G418. Correctly targeted embryonic stem cells were injected into C57BL6/J blastocysts to obtain chimeric mice and germline transmission. Heterozygous animals were intercrossed to obtain both mir-146a+/− and mir-146a3′F/3′F homozygotes.

**Animal studies**

Transgenic founder animals from each line were individually backcrossed three times to the C57BL6/J line (Jackson) and breeding colonies were separately maintained for each line, using heterozygotes for each mutant allele to produce paired homozygous mutant and wild-type control animals for each line. Age- and sex-matched mice from each of the two lines were used for all experiments (except where the use of sex-matched littermates is explicitly noted) such that homozygous mutants from each line were compared with an equivalent number of wild-type mice equally derived from the same two lines. No differences were noted in the phenotypic comparison of wild-type mice derived from breeding of mir-146a+/− heterozygotes and wild-type mice derived from breeding of mir-146a3′F/− heterozygotes in any experiments. All mouse procedures were approved by the Institutional Animal Care and Use Committees of the Baylor College of Medicine.

**Direct quantitation of WT and 3′F mir-146a**

Liver, thymus, kidney, lymph nodes, spleen, and whole bone marrow were excised from age- and sex-matched WT, mir-146a−deficient, and mir-146a3′F/3′F mice and physically disaggregated. Splenocytes were separated into B cells and CD4+ and CD8+ T cells via flow sorting using a FACSAria II (BD). Bone marrow cells were cultured for 1 wk in either IL-4 (10 ng/ml) + GM-CSF (30 ng/ml) or M-CSF (10 ng/ml) (all R&D Systems) for differentiation to either BMDCs or BMDCs, respectively. Differentiation was confirmed via flow cytometry using an LSRFortessa (BD). BMDCs and BMDCs were stimulated with 1 ng/ml LPS for 16 h. RNA was isolated from all samples using TRIzol LS reagent (Life Technologies) as per the manufacturer’s recommendations and quantified by spectrophotometry. Commercially obtained WT and 3′F mir-146a mimic direct quantitation standards were serially diluted in 1 μg/ml total RNA derived from Drosophila S2 cells. Reverse transcription was performed with the miScript II RT kit (QIAGEN) and qPCR was performed using the miScript SYBR Green PCR kit (QIAGEN), both according to the manufacturer’s instructions. Absolute miRNA quantification was calculated based on total input RNA for each biological sample. Primers specific for either the WT or the 3′F allele were used on all samples to ensure accurate genotyping and to rule out cross-primer.

**Luciferase reporter constructs and luciferase assay**

The 3′ UTRs of indicated genes were amplified from mouse genomic DNA via PCR and subcloned into an edited version of the pRL-TK-CX6X Renilla luciferase reporter plasmid (Addgene) via Xhol and Apal or Xhol and NotI sites added to the 5′ and 3′ ends of the PCR amplicons. Following restriction digestion, amplicons were ligated with the vector using the NEB Quick Ligation kit (NEB). Predicted mir-146a recognition sites within the UTRs were mutated as indicated in the text using the QuikChange II site-directed mutagenesis kit (Thermo Fisher Scientific). A more detailed catalog of oligonucleotide sequences is provided in Table S4.

HeLa cells were seeded in 24-well plates at a seeding density of 40,000 cells/well in DMEM supplemented with L-glutamine, penicillin, streptomycin, sodium pyruvate, Hepes (pH 7.4), nonessential amino acids, and β-mercaptoethanol (DMEM complete) and allowed to grow overnight. The following day, the cells were transfected with the pGL3 Firefly luciferase control plasmid (Promega), one of the experimental Renilla luciferase reporter plasmids, and varying concentrations of either the WT or 3′F mirRNA mimic duplexes (Sigma-Aldrich), along with a balanced amount of irrelevant anti–CXCR4 siRNA duplex (Sigma-Aldrich) to ensure that the final siRNA concentration in each well was always equal to 31.6 nM. Transfections were performed using the Lipofectamine 3000 transfection reagent system (Thermo Fisher Scientific) as per the manufacturer’s protocol. Luciferase assays were performed using the Dual-Luciferase Reporter Assay System (Promega) as per the manufacturer’s protocol on a Tecan M200 multimode reader using Tecan Magellan software (Tecan).

**Splenic immunophenotyping**

Spleens were excised, massed, photographed, and disaggregated. Red blood cells were lysed with ACK buffer, and the splenocyte suspensions were quantified before staining for the indicated markers. One million splenocytes were stained using a 1:100 dilution of fluorophore-tagged antibodies in 100 μl PBS + 1% FBS + 1 mM EDTA for 15 min at 4°C, after which the cells were washed three times. Cell viability was ensured by staining with NucBlue Live ReadyProbes Reagent (Thermo Fisher Scientific). Cells heat-killed at 65°C for 15 min were used as a control. Data were collected using a BD LSRFortessa flow cytometer and analyzed using Kaluza Analysis software V1.5 (Beckman Coulter). A detailed catalog of antibodies utilized is provided in Table S4.

**Histological analysis**

Livers and kidneys were excised to 70% ethanol and submitted to the Baylor College of Medicine Center for Comparative Medicine, where they were processed for hematoxylin and eosin staining via standard methods. Histological scoring was performed by a staff licensed veterinary pathologist blinded to the origin of the histological samples.
LPS challenges

For sublethal LPS challenges, mice were bled retro-orbitally and then stimulated with LPS at a dose of 1 mg/kg body mass. The mice were bled again at 2 h and 24 h post-injection. For lethal LPS challenges, the mice were retro-orbitally bled and then stimulated with LPS at a dose of 35 mg/kg body mass. The mice were bled again at 6 h post-injection. Serum was collected from the blood via differential centrifugation at 9,800 g for 7 min at 4°C. IL-6 was quantified via commercial ELISA kit (R&D Systems) following the manufacturer’s instructions. Other cytokines were quantified via Lumines assay (Thermo Fisher Scientific) by the Baylor College of Medicine Antibody-Based Proteomics core. For lethal LPS challenges, the mice were monitored hourly for morbidity beginning at 10 h post-injection.

CBC analysis

Blood was collected via retro-orbital bleeding into EDTA-coated collection vials (BD Pharmingen) and submitted to the Baylor College of Medicine Center for Comparative Medicine.

Isolation and culture of BMDMs

Bone marrow from 6- to 7-mo-old mice was flushed from femurs and tibias and plated at 85,000 cells/cm² on non—tissue culture—treated petri dishes. The cells were cultured for 8—9 d in DMEM complete medium supplemented with M-CSF (10 ng/ml), with a medium change at day 3. Successful differentiation was verified by surface staining of CD11b, CD11c, F4/80, CD86, and GR-1, followed by flow cytometry analysis as described above.

In vitro BMDM LPS challenges

BMDMs were stimulated with 1 ng/ml LPS for 24 h. To allow monitoring of cytokine production specifically within various windows following stimulation, cell culture medium was collected and replaced with medium supplemented with 1 ng/ml LPS every 8 h following stimulation. The culture medium was snap-frozen in liquid nitrogen and stored at −80°C. Analysis of IL-6 content in the medium was carried out via commercial ELISA kit (R&D Systems) according to the manufacturer’s instructions.

Immunoblots

BMDMs were lysed in RIPA buffer (10 mM Tris [pH 7.4], 150 mM NaCl, 1% Triton X-100, 0.1% SDS, 0.5% Na-Desoxycholate, 1 mM EDTA, and 1 mM DTT) containing protease and phosphatase inhibitors, snap-frozen, and stored at −80°C. Lysate protein concentration was quantified by the BCA protein assay. 40 μg of sample was diluted in 1× RIPA buffer, denatured in Laemmli buffer at 90°C for 10 min, and loaded into a 4—12% polyacrylamide gel. The gels were transferred to a PVDF membrane at 30 V for 1 h, blocked in 10% non-fat milk in TBST, and then stained overnight at 4°C with primary anti-IRAK1 (CST) and anti-GAPDH antibodies (EMD Millipore). The next day, the blots were washed three times with TBST, then incubated in HRP-conjugated secondary antibody for 30 min at room temperature. Blots were developed with ECL 2 solution (Thermo Fisher Scientific) and exposed to film. A detailed catalog of antibodies utilized is provided in Table S4.

RNA Seq

Total RNA was extracted using Trizol (Thermo Fisher Scientific) according to the commercial protocol. 100 ng of total RNA was used for rRNA depletion using Ribo-Zero Gold rRNA removal Kit (Illumina) and following a modified protocol for low inputs. Half of the rRNA-depleted RNA was used for library preparation using NEBNext Ultra II Directional RNA Library Prep Kit (New England Biolabs), followed by sequencing on a NextSeq500 instrument (Illumina). The raw reads were processed to remove the adapter sequences using cutadapt (v1.8.3) with the following arguments: --min-intronlen 70, --min-strandness R, --known-splicesite-infile. The raw read counts were obtained for each gene (Gencode’s comprehensive gene annotation release M15) using featureCounts (v1.5.3) with the following arguments: -s 2, -Q 50. For the miRNA targeting analysis, log₂ fold-change of predicted targets of miR-146 (TargetScanMouse v7.1) was compared with the log₂ fold-change of background set (mRNAs that do not contain miRNA target sites in their 3’ UTR). The significance of the difference was evaluated using Wilcoxon rank sum test. Small RNA sequencing was performed essentially as described (Wissink et al., 2016). Raw RNAseq data may be obtained via NCBI GEO accession number GSE125474.

Mass spectrometry

Mass spectrometric analysis was carried out on an Orbitrap Fusion LumosETD (Thermo Fisher Scientific) by the Mass Spectrometry Proteomics core at Baylor College of Medicine as described before (Saltzman et al., 2018). Briefly, BMDM lysates from 8- to 12-wk-old mice were denatured in ABC solution (50 mM ammonium bicarbonate + 1 mM CaCl₂), with subsequent snap-freeze/4°C thaw cycling and boil denaturing at 95°C. The proteins were digested with a 1:20 solution of 1 μg/μl trypsin:protein overnight at 37°C with shaking, and then with a 1:100 solution of 1 μg/μl trypsin:protein for 4 h. The peptides were extracted and measured using the Pierce Quantitative Colorimetric Peptide Assay (Cat. No. 23275; Thermo Fisher Scientific). Next, 50 μg of vacuum-dried peptides were re-dissolved in pH10 ABC buffer (10 mM ammonium bicarbonate, pH 10, adjusted with NH₄OH) and subjected to off-line microscaled reverse-phase separation on a microporpett tip layered with 6 mg of C18 matrix (Reprosii–Pur Basic C18, 3 μm, Dr. Maisch GmbH) on top of a C18 disk plug (EmporeTM C18, 3M). A total of 15 (2–30% ACN, 2% steps) fractions were obtained and combined into five pools for mass spectrometry sequencing (15F5R protocol with 02+12+22, 04+14+24, 06+16+26, 08+18+28, and 10+20+30% combinations). For each
run on the Lumos instrument, ~1 µg of peptide was loaded onto a 2-cm 100 µm ID pre-column and resolved on a 6-cm 150 µm ID column, both packed with 2-µm C18 beads (Repersil-Pur Basic C18, Cat. No. r119.b9.0003, Dr. Maisch GmbH). The gradient mobile phase was mixed from water (solution A) and 90% acetonitrile (solution B), both with 0.1% formic acid. A constant flow rate was maintained with 75-min linear gradient elutions. The Proteome Discoverer software suite (PD version 2.0.0.802; Thermo Fisher Scientific) was used to search the raw files with the Mascot search engine (v2.5.1, Matrix Science), validate peptides with Percolator (v2.05), and provide MS1 quantification through Area Detector Module. MS1 precursors in a 350–10,000 mass range were matched against the trypptic RefProtDB database digest with Mascot permitting up to of two missed cleavage sites (without cleavage before P), a precursor mass tolerance of 20 ppm, and a fragment mass tolerance of 0.5 D. The following dynamic modifications were allowed: acetyl (protein N-term), oxidation (M), carbamidomethyl (C), DeStreak (C), and Deamidated (NQ). For the Percolator module, the target strict and relaxed FDRs for PSMs were set at 0.01 and 0.05 (1 and 5%), respectively, gpGrouper was used to assemble peptide identifications into gene products, and the final proteome data matrix were filtered to proteins with at least two identifications within a treatment/genotype triplicate. Protein groups were quantified by iBAQ values and median-normalized. Raw mass spectrometry data may be obtained from the PRIDE partner repository under accession number PXD011413.

**Putative novel miR-146a target identification**

The difference in log2 fold-change of proteins upon LPS stimulation was calculated for WT and KO (and separately, WT and 3'F) BMDMs. The respective lists were sorted on this difference, and candidate targets were drawn from two pools of gene products with a Z score of >3 in the positive or negative direction from the median. Candidate novel targets were defined as those with a putative seed in the CDS or 3' UTR of the corresponding mRNA and characterized by a significant change (q < 0.05) in one or both of the genotypes compared. These candidates (along with negative controls characterized by differential expression but no predicted seed region) were amplified from murine genomic DNA via PCR and inserted into the pRL-TK-CX6X backbone, as described above.

**Statistics**

For comparisons between two groups, statistical significance was assessed using a two-tailed t test both with and without assumption of equal variance. Neither assumption impacted the level (or lack thereof) of significance depicted in any representation. Data are presented as mean ± SEM in pooled experiments and mean ± SD in representative individual experiments. Comparisons of different miRNA mimics across multiple concentrations were analyzed by two-way ANOVA. Survival curves were evaluated using Mantel–Cox log-rank test. In all cases, * denotes a P-value of <0.05 and ** denotes a P-value of <0.01. All statistical analyses were performed using GraphPad Prism v6.0.

**Supplementary Information**

Supplementary information is available at https://doi.org/10.26508/lsa.201800249.

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**Conflict of Interest Statement**

The authors declare that they have no conflict of interest.

**References**


