1 Supplemental experimental procedures

1.1 Measurement of inter-individual variability in gene expression

Source code and data files are available at:

http://www.igh.cnrs.fr/equip/Seitz/Pinzon_et_al_2016_data_and_code/Figure_1.tar.bz2.

Ten adult (3 month old) male, pathogen-free S/SPF C57BL/6J mice (from Charles River Laboratories; accustomed to our mouse facility for 9 days before the experiment) were sacrificed according to local regulations. Approval for these studies was obtained from the Ethics Committee on Animal Research of the Languedoc-Roussillon region (CE-LR-0505). 800 to 900 μ L blood was collected from each mouse and heparin-treated. Five blood samples were analyzed separately ("biological replicates") and the other five samples were pooled, then split into five "technical replicates". All ten samples were then treated identically in a double-blind manner.

Neutrophil RNA was isolated using the anti-mouse Ly6G PE antibody (BD Biosciences cat. # 551461), anti-PE magnetic beads (Miltenyi Biotec cat. # 130-048-801), MS columns (Miltenyi Biotec cat. # 130-042-201) and the QIAamp RNA extraction kit (Qiagen cat. #52304). Neutrophil purity (ranging from 71.5 to 99.5 %) was verified by flow cytometry. Neutrophil lysis and RNA extraction was performed on a QIAcube robot, for better sample-to-sample reproducibility. RNA was precipitated, DNase-treated, further purified using the RNeasy kit (Qiagen cat. # 74104), and handed to the IRB microarray facility (Montpellier, France).

cDNAs were labeled using the Affymetrix 3' IVT labeling kit with the two-cycle cDNA synthesis protocol, and hybridized on MG-430 PM array strips (Affymetrix cat. # 901570). Hybridization was performed at 45°C and washes were performed between 37.07 and 38.00°C, on an Affymetrix GeneChip station. Image acquisition was performed on a 3000 7G Affymetrix scanner. Intensity values were background-subtracted and normalized using the RMA method. Out of 196 predicted miR-223 targets according to TargetScan mouse v. 6.2, four are not probed by the array (genes 5031414D18Rik, Cdh12, Zfp839 and Fignl2). Baek et al. (2008)'s analysis, as well as ours, was thus restricted to the remaining 192 genes.

For each probeset on the array, we measured technical variability for probesets with similar signal intensities (*i.e.*: the 5 percentiles whose mean intensity is lower than the mean intensity of the probeset of interest, and the 5 percentiles whose mean intensity is higher). For each of the 5 technical replicates of each of these probesets, we computed the ratio: (replicate intensity)/(mean intensity across all 5 technical replicates). These ratios follow approximately a normal distribution (see middle panel of figure 1**B** in the main text for an example).

For each probeset, the probability of each possible underlying biological value for each individual mouse was inferred from such normal distribution (see right panel of figure 1**B** in the main text for an example). The *p*-values shown in figure 1**D** and in table S1 measure the probability that the median of the miR-223-guided fold-change in gene expression (taken from NCBI's GEO accession #GSE12001, described in Baek et al. (2008)) exceeds the measured inter-individual variability in gene expression. They were estimated by randomly picking 100,000 values in the distribution of possible underlying biological values for the most highly and lowly expressing mice, according to their estimated probability density. The *p*-value was defined as the frequency of random pairs whose difference is smaller than the median miR-223-guided fold-change. Classification into variable genes (p < 0.05) and tightly-regulated genes ($p \ge 0.05$) was based on the median *p*-value across probesets.

Using the Loess normalization method instead of RMA, even more genes appear to have a large interindividual variability: with a *p*-value cutoff of 0.01, only 6 genes (*Bai3*, *Fbxo8*, *Srp54a*, *Srp54b*, *Tgfbr3* and *Ube2q2*) had a median *p*-value above the cutoff (meaning that their inter-individual variability appears to be smaller than miR-223-guided repression).

Because neutrophil purity is not identical across individual mouse samples (see Supplemental Table S1), it is formally possible that contaminating cells contribute variable amounts of mRNA from genes that are mostly expressed in some non-neutrophil cells. Such contamination, which is variable across samples, could generate arbitrarily high artifactual variability. In order to control for this confounding effect, we scanned the probesets on the array to identify which sample exhibits the highest expression level. If indeed variable amounts of contaminating cells generate artifactual variability for non-neutrophilspecific genes, then one would expect the least pure neutrophil preparations to exhibit anomalously high expression levels for these non-neutrophil-specific genes. We thus scanned all 45,141 probesets on the microarray and recorded which of the 5 individual mouse samples exhibits the highest expression level (see Supplemental Table S1): probe level repartition does not seem to depend on neutrophil purity, ruling out the possibility that our observed inter-individual variability is due to variable contamination.

1.2Comparison of dose-sensitivity predictors with miRNA binding site conservation

Source code and data files are available at: http://www.igh.cnrs.fr/equip/Seitz/Pinzon_et_al_2016_data_and_code/Figure_2.tar.bz2.

The list of known human haplo-insufficient genes was taken from Dang et al. (2008). TargetScan 7.0's aggregate P_{CT} (probability of conserved targeting) scores were downloaded from http://www. targetscan.org/vert_70/vert_70_data_download/Conserved_Family_Info.txt.zip and http://www. targetscan.org/vert_70/vert_70_data_download/Nonconserved_Family_Info.txt.zip (see Friedman et al. (2009) for the description of the aggregate P_{CT}). For each gene, we considered the miRNA family with the highest aggregate P_{CT} (*i.e.*: miRNAs with the most conserved interaction to that gene). Every known haplo-insufficient gene described by Dang et al. (2008) is a predicted target for human miR-NAs according to TargetScan 7.0. Among TargetScan-predicted miRNA targets, the remaining 19,066 genes are not known as haplo-insufficient according to Dang et al. (2008): they were used as a control in Figure 2A.

The probability of human genes for being haplo-insufficient was calculated by Huang et al. (2010) for 12,218 genes. 9,520 of these exhibit conserved miRNA binding sites according to TargetScan 7.0. For each gene, we considered the miRNA family with the highest aggregate P_{CT} .

1.3Absolute quantification of miRNAs and their targets during C2C12 cell differentiation

Source code and data files are available at:

http://www.igh.cnrs.fr/equip/Seitz/Pinzon_et_al_2016_data_and_code/Figure_3.tar.bz2 and: http://www.igh.cnrs.fr/equip/Seitz/Pinzon_et_al_2016_data_and_code/Figure_4.tar.bz2

C2C12 cells were ordered freshly from ATCC (cat. #CRL-1772) then always kept sub-confluent. Differentiation was induced as described in Sweetman et al. (2008) once cells reached confluency.

1.3.1**RIP-based target identification**

Cells were transfected with 20 nM antisense oligonucleotides using Lipofectamine 2000 (Invitrogen) 51 hours after differentiation induction (3 replicates per transfection; seven confluent 150 mm diameter Petri dishes per replicate). 24 hours after antisense oligonucleotide transfection, cells were rinsed with cold PBS then cross-linked at 0 °C with 150 mJ.cm⁻² 254 nm U.V. light (through 12 ml PBS per 150 mm diameter Petri dish). Cells were collected immediately, cell suspension was divided into two equal parts, pelletted and flash-frozen in liquid nitrogen. One Petri dish out of seven was not cross-linked, but collected by trypsination and analyzed by flow cytometry for transfection efficiency quantification.

 $150 \,\mu\text{L}$ Dynabeads protein G suspension (Novex; $30 \,\text{mg.mL}^{-1}$) were washed 3 times in $1 \,\text{mL}$ PBS + 0.02%Tween 20, resuspended in 300 μ L PBS + 0.02% Tween 20, then incubated overnight at 4°C under gentle agitation with 3 μ L anti-mammalian Ago 2A8 ascites (a kind gift of Prof. Z. Mourelatos, University of Pennsylvania; $\approx 15 \ \mu g.\mu L^{-1}$). Beads were then washed 3 times in 1 mL PBS + 0.02% Tween 20. Immunoprecipitation was performed on one of the two frozen cell pellets for each replicate. Cells were disrupted in 3 mL lysis buffer¹ per cell pellet, then incubated on ice for 10 min. Lysates were DNasetreated², then ultra-centrifuged 20 min at 35,000 g at 4 °C. The supernatant (after sampling 350 μ L for input control) was incubated with 2A8-bound Dynabeads for 1h30 at 4°C under gentle agitation. $350 \ \mu L$ of supernatant were kept for control, and beads were successively washed with 2 mL of the following buffers at 4°C:

- lysis buffer¹;
- high salt buffer³
- high stringency buffer⁴
- low salt buffer⁵

¹PBS with 1% Empigen, 40 U.mL⁻¹ RNasin (Promega), Protease inhibitor cocktail (Roche) (1 tablet per 10 mL). ²36 μ L RQ1 DNase (Promega; 1 U. μ L⁻¹) for 3 mL of lysate, incubated 8 min at 37 °C. ³PBS with 1% Empigen, supplemented with 50 g.L⁻¹ NaCl.

⁴15 mM Tris-HCl (pH=7.5), 5 mM EDTA (pH=8.0), 2.5 mM EGTA (pH=8.0), 1% Empigen, 120 mM NaCl, 25 mM KCl.

⁵15 mM Tris-HCl (pH=7.5), 5 mM EDTA (pH=8.0).

• PNK buffer⁶

then optionally, 33 μ L of bead suspension in final wash were kept for Western blotting control. Washed beads were resuspended in 100 μ L proteinase K buffer⁷ and 2.5 μ L proteinase K (New England Biolabs; 20 μ g. μ L⁻¹), incubated at 37°C for 30 min then extracted in 100 μ L phenol/chloroform/isoamyl alcohol (25/24/1) and RNA was precipitated with 50 μ g glycogen, 10 μ L NaAcO (3 M, pH=5.2) and 300 μ L ethanol. Recovered RNA quantities ranged from 184 to 289 ng.

RNA samples were handed to Beijing Genomics Institute for library preparation (poly(A)-independent protocol, fragmenting RNAs prior to reverse-transcription; 49 nt reads, single-end sequencing; adapter sequence: 5'-AGATCGGAAGAGCACACGTCTGAACTCCAGTCAC-3'). After adapter trimming, reads shorter than 20 nt (*i.e.*: 0.14% to 1.96% of each library) were discarded.

Differential mRNA abundance between anti-miR-1a/miR-206 and anti- \emptyset libraries, and between anti-miR-133 and anti- \emptyset libraries was identified using HTseq (http://www-huber.embl.de/HTSeq/doc/overview.html) and EdgeR (https://bioconductor.org/packages/release/bioc/html/edgeR.html) (normalizing read counts with the TMM method). "Experimentally identified targets" were defined as mRNAs which are significantly (adjusted *p*-value < 0.05) less abundant in anti-miRNA transfection libraries than in anti- \emptyset libraries.

1.3.2 Absolute RNA quantification

At each time point (every 24 h, from 0 to 6 days after differentiation induction), cells from three indepent Petri dishes (150 mm diameter) were collected in 10 mL DPBS. 500 μ L of the cell suspension was mixed with 4 μ L Thiazole Orange, 10 μ L of suspension were then monitored by microscopy to measure cell size. The remaining 494 μ L were mixed with 50 μ L Counting Beads (BD Biosciences cat. #349480, 1,043 beads. μ L⁻¹) and analyzed by flow cytometry to determine cell number, following the manufacturer's intructions.

The remaining 9.5 mL of initial cell suspension was used for RNA extraction with Trizol (measuring aqueous phase loss after centrifugation by weighting). 60 to 300 μ g RNA was obtained for each replicate.

20 μ g total RNA were used for Northern blotting for miRNA quantification. A mix of 27 *in vitro* transcribed and polyadenylated spike-ins (not matching the mouse genome) was prepared and verified by Northern blot. Transcript concentration in the mix span a 100-fold concentration range. Spike-in mix was added to $\approx 10 \ \mu$ g total RNA from day 0, 3 and 6 of differentiation then handed to the Beijing Genomics Institute for RNA-seq library preparation and sequencing using the poly(A)-dependent protocol.

RNA abundance was calculated from RNA-seq data using the TopHat v.2.0.10 (http://ccb.jhu.edu/ software/tophat/index.shtml) and Cufflinks v.2.1.1 (http://cufflinks.cbcb.umd.edu/) programs using the -b and -G options.

1.3.3 Tmsb4x miR-1a/miR-206 site mutagenesis

Homology donor sequences were prepared by PCR on C2C12 genomic DNA, with the left homology arm containing either the wild-type (CAUUCC) or a mutated (UCCAUC) seed match for miR-1a/miR-206 in the Tmsb4x 3' UTR (nt 30 to 35 of the UTR) (see Figure 4A for a schematic map). The left homology arm is 986 bp long and the right homology arm is 861 bp long.

Undifferentiated C2C12 cells were co-transfected using Lipofectamine 2000 with:

- sgRNA-expressing plasmid: pmU6-gRNA (Addgene #53187), see section 1.6 for insert oligo sequences;
- High-fidelity-Cas9-expressing plasmid: VP12 (Addgene #72247);
- Homology donor plasmid: pmirGLO (Promega cat.#E1330) containing wt or miR-1a/miR-206 site mutant left homology arm (cloned between BglII and MluI sites) and right homology arm (cloned between ClaI and BamHI sites). In both constructs, two miR-1a/miR-206 binding sites were cloned in the pmirGLO multiple cloning site (the miR-1a/miR-206 sites from Tpm4 and Tppp 3' UTRs, each embedded in 94 nt of genomic context; we chose these two genes among our list of experimentally-identified targets [for Tpm4] and among TargetScan-predicted targets that we did not detect experimentally [for Tppp]), to make the luc2 reporter sensitive to miR-1a/miR-206 activity. That plasmid also contains a fusion gene between Renilla luciferase and the neomycin

⁶20 mM Tris-HCl (pH=7.5), 10 mM MgCl₂, 0.2% Tween 20.

⁷100 mM Tris-HCl (pH=7.5), 50 mM NaCl, 10 mM EDTA (pH=8.0).

resistance gene ("hRluc-neo^R fusion") which provides both a normalizing luciferase activity and an antibiotics resistance gene for stable cell line selection.

Stable polyclonal lines were selected by growing subconfluently in 200 to 600 μ g.mL⁻¹ G418 for 8 weeks. Five independently-transfected wild-type lines and four independently-transfected mutant lines were finally isolated and used for further analysis. The frequency of wild-type and mutant alleles in each line was verified both by qPCR (see Supplemental Figure S5) and low-throughput sequencing of their genomic *Tmsb4x* 3' UTR (not shown). The nine cell lines were differentiated for 6 days using the protocol described by Sweetman et al. (2008) then luc2 and hRluc activities were measured using the Dual luciferase reporter assay system (Promega cat.#E1910).

1.4 Correlation analysis between mRNA abundance and miRNA binding site conservation

Source code and data files are available at:

http://www.igh.cnrs.fr/equip/Seitz/Pinzon_et_al_2016_data_and_code/Figure_5.tar.bz2

For each murine miRNA family (as defined in miRBase v17), conservation of its interaction with predicted targets was estimated by the aggregate probability of conserved targeting of its miRNA binding sites (Friedman et al., 2009). For each probeset on the array, mRNA abundance was defined as the mean array signal across available biological replicates. Kendall's τ correlation coefficient was then calculated between these two datasets, for every miRNA family and every tissue.

1.5 Comparison between seed match conservation and seed conservation

Source code and data files are available at: http://www.igh.cnrs.fr/equip/Seitz/Pinzon_et_al_2016_data_and_code/Figure_6.tar.bz2.

miRNA seed match coordinates were extracted from the 100-species whole genome alignment centered on the human genome (http://hgdownload.cse.ucsc.edu/goldenPath/hg19/multiz100way/maf/). 3' UTR coordinates were retrieved from the UCSC Genome Browser RefSeq gene database (http:// hgdownload.soe.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz). For each human 3' UTR seed match, the list of vertebrate species having that seed match conserved in the genome alignment was extracted.

For vertebrate species listed in miRBase (release 21), the existence of each seed in the miRNA repertoire of a given species was extracted from miRBase. It is likely that some miRNAs have escaped experimental identification in some species: in order to avoid such false negatives, and also to determine whether species not recorded in miRBase express miRNAs with a given seed, we searched these species' genomes with HMMer for potential orthologous pre-miRNAs of known vertebrate miRNAs with the seed of interest:

- 1. known vertebrate miRNAs with that seed were collected, and their pre-miRNAs were aligned with blast;
- HMMer profiles were built for each group of mutually similar pre-miRNA sequences (pairwise blast E-value<0.001); if the group contains only 1 pre-miRNA, no HMMer profile could be built: homology search was then performed with blast;
- 3. each genome outside the clade of interest was searched for homologous loci using **nhmmer** with these profiles (when possible), or with **blast** otherwise;
- 4. each homologous locus (with an E-value < 0.001) was folded using RNAsubopt to predict transcript secondary structure; loci predicted to be transcribed in unbranched hairpins (whose predicted ΔG was at most 75% of the most stable structure's ΔG) were selected;
- 5. potential pre-miRNA orthologs were considered if their miRNA seed is identical to the seed being searched.

Seeds for which we found potential pre-miRNA orthologs in a genome outside the clade of interest were discarded.

For comparison with miRNA prediction programs, predicted miRNA binding sites were downloaded from:

- http://cbio.mskcc.org/microrna_data/human_predictions_S_C_aug2010.txt.gz and http: //cbio.mskcc.org/microrna_data/human_predictions_S_0_aug2010.txt.gz (miRanda aug2010);
- http://www.targetscan.org/vert_70/vert_70_data_download/Conserved_Site_Context_Scores. txt.zip and http://www.targetscan.org/vert_70/vert_70_data_download/Nonconserved_Site_ Context_Scores.txt.zip (TargetScan 7.0);
- http://diana.imis.athena-innovation.gr/DianaTools/data/microT_CDS_data.tar.gz (microT 5.0);
- and http://dorina.mdc-berlin.de/ (selecting the PicTar2 mammalian miRNA:targets in 3' UTRs) for PicTar2.

Prediction programs do not have the exact same requirements for target predictions, but they all tend to favor miRNA binding sites with extensive pairing to the seed. In order to make every analysis comparable, they were restricted to perfect seed matches in 3´ UTRs, excluding seed matches that overlap exon-exon junctions.

1.6 Oligonucleotides used in this study

RNA and 2-O-methyl oligos:

	Description	Sequence $(5 \text{ 'to } 3 \text{ '})$				
As	anti-miR-1a	$\verb"Cy5-AmCmGmAmUmAmCmAmUmAmCmUmUmCmUmUmAmCmAmUmUmCmCmAmAmCmGm-ddC" \\$				
iRN	anti-miR-206	${\tt y5-AmCmGmCmCmAmCmAmCmAmCmUmUmCmCmUmUmAmCmAmUmUmCmCmAmAmCmGm-ddC}$				
ti-m	anti-miR-133	$\verb"Cy5-AmCmGmCmAmGmCmUmGmGmUmUmGmAmAmGmGmGmGmAmCmCmAmAmAmAmCmGm-ddC""$				
an	anti- \varnothing	$\verb"Cy5-AmCmGmAmUmAmAmCmGmUmAmCmGmUmAmCmGmUmAmCmGmUmAmCmGmUmAmCm-ddC" \\$				
As	miR-1a	P-UGGAAUGUAAAGAAGUAUGUAU				
RN	miR-206	P-UGGAAUGUAAGGAAGUGUGUGG				
В	miR-133	P-UUUGGUCCCCUUCAACCAGCUG				

DNA oligos:

	Description	Sequence $(5 \text{ to } 3)$
sis	Left homology arm cloning (forward)	ttacgcgtTGATGAATATGGCCTGCAAA
te ne	Left homology arm cloning (reverse)	ttagatctCTACCCCTTCATTCCACAGC
utag	Right homology arm cloning (forward)	ttatcgatGCTGTGGAATGAAGGGGTAG
r H	Right homology arm cloning (reverse)	ttggatccAAAGGCCTGTTCACTGATGG
ms b4	sgRNA cloning (top strand)	P-TTGTTTGCAAGCTGGCGAATCGTAATG
T	sgRNA cloning (bottom strand)	P-AAACCATTACGATTCGCCAGCTTGCAA
CR	Forward primer for qPCR (common to wt and mutant allele)	AGCGAGGCTGCTATGTGTCT
^q P	Reverse primer for qPCR on wt allele	AAGGCAATGCTCGTGGAAT
	Reverse primer for qPCR on mutant allele	AAGGCAATGCTCGTGATGG
rter	Tpm4 miR-1a/miR-206 site cloning (forward)	aagagctcTGGCTTTGTAGTTTTCCTTTCTC
repo	Tpm4 miR-1a/miR-206 site cloning (reverse)	aactcgagTCAGGACTGTAAACTTGAGTTGG
	Tppp miR-1a/miR-206 site cloning (forward)	aactcgagCACTATAGGTGGCAGGCACA
lim	Tppp miR-1a/miR-206 site cloning (reverse)	aatctagaGCCTAGTGGAGGTGCATTCT

Cy5: Cyanine5; ddC: dideoxy C; Nm: 2´-O-methylated nucleotide; P: phosphate. Lower case letters: extragenomic sequences, for restriction site addition.

2 Supplemental figures



Figure S1: Inter-individual variability in gene expression in mouse neutrophils. Measured interindividual variability in gene expression for the most accurately quantified genes (mean of the 5 individual measured $\log_2(\text{signal}) > 2 \times \text{standard}$ deviation of $\log_2(\text{signal})$ of probesets with similar microarray signal). Each point represents one gene. The *y*-axis displays the maximal measured inter-individual variability (signal in the most strongly expressing mouse / signal in the least strongly expressing mouse). miR-223 targets were predicted using TargetScan mouse (v. 6.2). For graphical clarity, only the genes for which variability is less than 10-fold are represented (that is: 177 predicted targets and 20,542 other genes). Variability for the remaining 12 predicted targets and 947 other genes reaches 46-fold and 197-fold, respectively (not shown).



Figure S2: Uncoordinated expression of predicted miR-223 targets among individual mice. For each predicted miR-223 target, we ranked the 5 mice according to that gene's expression. For genes matched by several probesets on the array, we considered the mean of the ranks for every probeset. Each point on the plot represents a predicted miR-223 target. Rank distribution is rather uniform across the 5 mice, indicating that predicted targets are not coordinately up- or down-regulated from one individual to the next.



Figure S3: Anti-miRNA oligonucleotide transfection control. Flow cytometry was performed with the same acquisition settings for every sample. Recorded events with a high FSC (forward-scattering light; FSC cutoff=850 with our settings) were flagged as "cells" and selected for fluorescence analysis. Percentages of cells with a Cy5 fluorescence higher or lower than our cutoff (100 with our settings) are indicated in red on each histogram (rounded to the closest integer).



Figure S 4: A. In vitro transcribed spike-ins in calibrated RNA-seq. In each of the 9 RNAseq libraries, 27 polyadenylated *in vitro* transcripts were introduced in controlled amounts for internal calibration (each *in vitro* transcript is represented by a circle). For each transcript, between 3×10^{-17} and 3×10^{-15} mol were added in each RNA sample (RNA samples were prepared from $\approx 243,000$ to $\approx 780,000$ cells each). Linear regression of spike-in data, forcing a null *y*-intercept, then allowed us to convert *fpkm* RNA-seq results into numbers of molecules per cell, with a precision of ≈ 2 -fold (the linear regression line is shown as a plain black line). Please note the data is not log-transformed. **B. C2C12 differentiation control.** Unsupervised clustering of RNA-seq data (for mRNAs with more than 100 fpkm in at least one library) reveals the similarity of biological replicates at each differentiation (rows). Distance between features was measured by (1 - Kendall's τ correlation coefficient) and clustering was performed using the McQuitty method. Very similar results were obtained with alternative distance definitions and clustering procedures (not shown). *Id1* and *Id3* are known markers of the undifferentiated state of C2C12 cells (Jen et al., 1992; Mohamed et al., 2013) while *Mylpf*, *Myog* and *Casq2* are specifically expressed upon differentiation (Arai et al., 1992; Yuasa et al., 2009; Wu et al., 2014).



Figure S5: **qPCR-based quantification of wt and mutant** Tsmb4x alleles in the established **polyclonal lines.** qPCR reactions were performed on 100 ng genomic DNA, supplemented with either water or 4×10^{-13} g of various plasmid mixes. Plasmids were 4 kb long and contained either the Tmsb4x 3' UTR miR-1a/miR-206 seed match or the mutant version (CAUUCC—)UCCAUC); they were mixed in various proportions, with 0, 25, 50, 75 or 100% of either version. Each qPCR was performed as two technical replicates, except for lines "wt3" and "wt4".



Figure S6: The most highly expressed genes tend to bear the most conserved miRNA binding sites. Same conventions as in Figure 5A in the main text.



Figure S 7: Identification of clade-specific miRNA seeds. 100 vertebrate species are used in the human-centered whole genome alignment (Rosenbloom et al., 2015); species listed in miRBase release 21 (Kozomara and Griffiths-Jones, 2014) are shown in red. For each clade containing *Homo sapiens* we identified miRNA seeds that are present in at least 75% of the species in the clade, while being absent in every species outside the clade. It is likely that some miRNAs have escaped experimental identification in some species: in order to avoid such false negatives, these species' genomes were searched for potential orthologous pre-miRNAs of known vertebrate miRNAs with that seed. We excluded every seed for which potential pre-miRNA orthologs were found in genomes outside the clade of interest. Only the clades for which ≥ 10 clade-specific seeds could be identified were analyzed in Figure 6.

3 Supplemental tables

			Probesets with	Probesets with
Sample	Neutrophil purity (%)	Usage	highest signal	lowest signal
			among individuals	among individuals
1	90.5	Individual mouse	5,631	4,987
2	88.8	Pooled blood	_	_
3	85.0	Individual mouse	3,296	7,009
4	91.6	Pooled blood	_	_
5	71.5	Individual mouse	9,979	8,544
6	92.7	Pooled blood	_	_
7	89.8	Pooled blood	_	_
8	88.1	Pooled blood	_	_
9	99.5	Individual mouse	5,562	12,606
10	88.4	Individual mouse	$20,\!673$	11,995

Table S1: Neutrophil purity in the analyzed samples. Purity was assessed by flow cytometry using APC anti-mouse Ly6G antibody. "Individual mice" were used for measurement of inter-individual variability; "pooled blood" was used for measurement of technical inaccuracies. For "individual mouse" samples, we recorded the number of probesets for which the highest or lowest signal among individual mouse samples was found in that sample for quality control (see section 1.1).

	Tightly-regu	lated genes	Variable genes			
	Arfip1	Cbx5	A930038C07Rik	Abhd13	Ankrd17	Atp1b1
	Rasa1	Sept8	Atp2b1	F3	Fa2h	Fam 176a
	Srsf12		Foxp1	Hlf	Igf1r	Itgb1
Tarbase-supported			Mbnl1	Mef2c	Mga	Mtpn
interactions			Myh10	Myst3	Nfia	Nfib
			Otud4	Pds5b	Phf20l1	Ptbp2
			Rap2a	Rhob	Slc37a3	Sox 6
			Srgap3	Stk39	Y pel 1	
	Aco1	A cvr2a	Acsl3	Adcy7	Aebp2	Armc1
	Anks1b	Arpp19	Armcx1	Atp 7a	Atxn1	Bai3
	Brpf3	Ctsl	Cd2ap	Cdk17	Celf2	Cep68
	Fam 199x	Fam 46a	Cnot2	Cntln	Cops2	Cpne4
	Fbxo25	Fbxo8	Crim1	Csnk1g1	D19Wsu162e	Ddit4
	Fgfr2	Frmd4a	Dennd5b	Dlc1	Dusp2	E330021D16Rik
	Gpr22	Lelp1	Ebf3	Eif1ad	Eif2c3	Eif5b
	Mospd1	Nlrp3	Elf5	Elk1	Ept1	Fam 120c
	Orc4	Pik3c2a	Fam 168a	Fam5c	Fat1	Fbxw7
	Plce1	Rc3h1	Flrt1	Galnt7	Gna13	Gpm6b
	Rcn2	Scn1a	Gpr155	Hhex	Hsp90b1	Il6st
	Sh3pxd2b	Slc23a2	Inpp4a	Inpp5b	Jmy	Kif21b
	Slc35f1	Slc39a1	Lass6	Lif	Lmo2	Mafb
	Smarcd1	Srp54a	Mll3	Mmp16	Mon2	Mpz
Other interactions	Srp54b	Tet3	Msi2	Mtap4	Naa50	Nfat5
	Tgfbr3	Tmem 170	Nlgn2	Nutf2	Olfm1	Pde4d
	Tmprss 11e	Ube2q2	Phactr4	Phip	Pkn2	Pknox1
	Ulk2	Zcchc14	Plagl 2	Plekhh1	Prpf39	Rab22a
	Zxdc		Rab8b	Rabgap1	Rabl2	Rad17
			Ralgps 2	Rbm16	Rbm20	Rbpj
			Rnf34	Rnpc3	Rorb	Rps6kb1
			Rras2	Scn2a1	Scn3a	Sept6
			Sgms2	Slain2	Slc24a2	Slc4a4
			Slc8a1	Smoc1	Smurf2	Sox 11
			Sp3	Spata 13	Srp54c	Srpk2
			Styx	Syncrip	Tmem 229a	Tmem 47
			Tmem64	Tmtc2	Tnrc6b	Treml2
			Trps1	Tshz3	Ube2a	Uqcc
			Vamp2	Vav3	Wdr62	Wwtr1
			Zfhx3	Zfp238	Zfx	

Table S 2: Tightly-regulated genes and variable genes among predicted miR-223 targets. "Tightly-regulated genes" are the predicted miR-223 targets whose inter-individual variations in expression are lower than miR-223-guided repression (p < 0.05, with *p*-value defined as in Figure 1C). "Variable genes" are the other predicted targets. TarBase data was downloaded from http://diana.imis.athena-innovation.gr/DianaTools/index.php?r=tarbase/index. Among the 42 tightly-regulated genes, 5 are listed in TarBase; among the 150 variable genes, 31 are listed in TarBase (proportions not significantly different: Fisher's exact test *p*-value=0.37).

Differentiation	Replicate	Number	Genome	Transcriptome	Number of	Median abun-
day	name	of reads	matching	matching	detected	dance of detected
			reads	reads	mRNAs	mRNAs (fpkm)
0	1	343,336,245	336,133,200	326,332,352	17,481	3.46836
0	2	$329,\!549,\!176$	323,584,543	313,815,004	$17,\!356$	3.29692
0	3	339,119,201	332,961,581	$322,\!807,\!963$	17,475	3.25078
3	1	325,133,660	319,030,091	314,639,678	17,898	4.121915
3	2	$272,\!084,\!347$	267,228,824	$264,\!513,\!001$	17,785	4.19869
3	3	$319,\!687,\!481$	313,691,691	309,703,151	17,846	4.162125
6	1	313,253,641	307,337,296	303,106,144	18,061	4.09468
6	2	$339,\!291,\!429$	332,544,825	327,073,727	18,059	4.07273
6	3	$298,\!424,\!803$	$292,\!694,\!151$	$288,\!940,\!732$	18,115	4.1594

Table S3: **RNA-seq statistics.** The number of transcriptome-matching reads was rounded from Cufflinks' "normalized map mass" for each library. "Detected mRNAs" are the murine mRNAs for which read abundance > 0 fpkm.

NCBI's GEO sample	NCBI's GEO series	Organ or tissue
GSM461846	GSE18551	12 month striatum
GSM461847		
GSM461848		
GSM461849		
GSM461850	GSE18551	24 month striatum
GSM461852		
GSM461855		
GSM461859		
GSM252093	GSE9954	adipose tissue
GSM252094		
GSM252095		
GSM252086	GSE9954	adrenal gland
GSM252087		
GSM252088		
GSM252089	GSE9954	bone marrow
GSM252090		
GSM252091		
GSM252092		
GSM252077	GSE9954	brain
GSM252078		
GSM252079		
GSM149511	GSE6514	cerebral cortex
GSM149512		
GSM149513		
GSM149514	-	
GSM149515	-	
GSM252064	GSE9954	diaphragm
GSM252065		
GSM252066		
GSM200700	GSE8091	E11.5 embryonic head
GSM200701		, , , , , , , , , , , , , , , , , , ,
GSM200703		
GSM200704		
GSM243346	GSE9629	E12.5 embryonic kidney
GSM243347		
GSM243348		
GSM200706	GSE8091	E13.5 embryonic head
GSM200707		-
GSM200708		
GSM200709		
GSM200711		
GSM200713		
GSM200695	GSE8091	E9.5 embryonic head
GSM200696	1	
GSM200697	1	
GSM200698		
GSM200699		
GSM200716		
GSM252122	GSE9954	ES cells
GSM252123		
GSM252124		

(to	be	continued)
1		

NCBI's GEO sample	NCBI's GEO series	Organ or tissue
GSM252119	GSE9954	eye
GSM252120		
GSM252121		
GSM252131	GSE9954	fetus
GSM252132		
GSM252133		
GSM252113	GSE9954	heart
GSM252114		
GSM252115		
GSM149576	GSE6514	hypothalamus
GSM149577		
GSM149578		
GSM149599		
GSM149600		
GSM252083	GSE9954	kidney
GSM252084		
GSM252085		
GSM228786	GSE9012	liver
GSM228787		
GSM228788		
GSM228789		
GSM228790		
GSM252074	GSE9954	
GSM252075		
GSM252076		
GSM252080	GSE9954	lung
GSM252081		_
GSM252082		
GSM426400	GSE17812	memory P14 T cells
GSM426401		Ť
GSM252070	GSE9954	muscle
GSM252071		
GSM252072		
GSM252073		
GSM94741	GSE4142	naïve B cells
GSM94744		
GSM94745		
GSM252128	GSE9954	ovary
GSM252129		v
GSM252130		
GSM252096	GSE9954	pituitary gland
GSM252097		- • •
GSM252098		
GSM252099		
GSM252100		
GSM252125	GSE9954	placenta
GSM252126		-
GSM252127		
L		

(to be continued)

NCBI's GEO sample	NCBI's GEO series	Organ or tissue
GSM458136	GSE18358	renal glomeruli
GSM458140		
GSM458141		
GSM458142		
GSM458143		
GSM252101	GSE9954	salivary gland
GSM252102		
GSM252103		
GSM252104	GSE9954	seminal vesicle
GSM252105		
GSM252106		
GSM252116	GSE9954	small intestine
GSM252117		
GSM252118		
GSM261510	GSE10347	soleus muscle
GSM261511		
GSM261512		
GSM252067	GSE9954	spleen
GSM252068		
GSM252069		
GSM61837	GSE2826	splenic B cells
GSM61838		
GSM252110	GSE9954	testis
GSM252111		
GSM252112		
GSM252107	GSE9954	thymus
GSM252108		
GSM252109		
GSM261516	GSE10347	tibialis anterior muscle
GSM261517		
GSM261518		

 $Table \ S4: \ {\bf Transcriptomic\ datasets\ used\ for\ the\ analysis\ of\ correlation\ between\ mRNA\ abundance\ and\ miRNA\ binding\ site\ conservation.$

		1			
		Predicted sites in 3´ UTRs			
Target prediction	Total predicted	Overlapping Not on exon-exon junctions			
program	sites	exon-exon	No perfect	With perfect	
		junctions	seed match	seed match	
microT 5.0	32,400,362	444,545	nt 2–7: 15,322,673	nt 2–7: 12,240,601	
			nt 2–8: 23,309,068	nt 2–8: 4,254,206	
miRanda aug2010	4,417,884	0	nt 2–7: 1,266,616	nt 2–7: 3,151,268	
			nt 2–8: 1,966,976	nt 2–8: 2,450,908	
PicTar2	1,287,398	0	nt 2–7: 479,513	nt 2–7: 550,906	
			nt 2–8: 754,240	nt 2–8: 276,179	
TargetScan 7.0	14,542,205	21,157	nt 2–7: 496,637	nt 2–7: 9,473,770	
			nt 2–8: 4,328,678	nt 2–8: 5,641,729	

Table S 5: Predicted miRNA binding sites characteristics. Origin of data: http: //diana.imis.athena-innovation.gr/DianaTools/data/microT_CDS_data.tar.gz for microT 5.0 predictions; http://cbio.mskcc.org/microrna_data/human_predictions_S_C_aug2010.txt.gz and http://cbio.mskcc.org/microrna_data/human_predictions_S_0_aug2010.txt.gz for mi-Randa aug2010 predictions; http://dorina.mdc-berlin.de/ for PicTar2 predictions; http://www. targetscan.org/vert_70/vert_70_data_download/Conserved_Site_Context_Scores.txt.zip and http://www.targetscan.org/vert_70/vert_70_data_download/Nonconserved_Site_ Context_Scores.txt.zip for TargetScan 7.0. Predicted binding site location was compared to 3' UTR exon coordinates from the UCSC Genome Browser RefSeq gene data file (http://hgdownload.soe.ucsc.edu/goldenPath/hg19/database/refGene.txt.gz).

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