RNAi-mediated knockdown of mouse melanocortin-4 receptor in vitro and in vivo, using an siRNA expression construct based on the mir-187 precursor

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Abstract: RNA interference (RNAi) is a powerful tool for the study of gene function in mammalian systems, including transgenic mice. Here, we report a gene knockdown system based on the human mir-187 precursor. We introduced small interfering RNA (siRNA) sequences against the mouse melanocortin-4 receptor (mMC4R) to alter the targeting of miR-187. The siRNA-expressing cassette was placed under the control of the cytomegalovirus (CMV) early enhancer/chicken β-actin promoter. In vitro, the construct efficiently knocked down the gene expression of a co-transfected mMC4R-expression vector in cultured mammalian cells. Using this construct, we generated a transgenic mouse line which exhibited partial but significant knockdown of mMC4R mRNA in various brain regions. Northern blot analysis detected transgenic expression of mMC4r siRNA in these regions. Furthermore, the transgenic mice fed a normal diet ate 9% more and were 30% heavier than wild-type sibs. They also developed hyperinsulinemia and fatty liver as do mMC4r knockout mice. We determined that this siRNA expression construct based on mir-187 is a practical and useful tool for gene functional studies in vitro as well as in vivo.

Key words: mir-187, mMC4R, RNAi, Transgenic mice

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RNA interference (RNAi) is a sequence-specific gene-silencing mechanism widely conserved among eukaryotic cells [19]. RNAi is triggered in mammalian cells by small interfering RNAs (siRNAs) that are 21 nucleotides (nt) long and contain 2 nt 3′ overhangs.

Animal cells contain numerous endogenous ~22 nt RNAs known as microRNAs (miRNAs) [16] that can cause mRNA degradation and/or translational inhibition when bound to partially complementary sites in the 3′ untranslated region (3′-UTR) of mRNAs [32, 34]. Natural miRNAs are transcribed by RNA polymerase II (pol II) as long, primary, 5′-capped, and polyadenylated miRNAs (pri-miRNAs) [3, 17]. The pri-miRNA is cropped first to release a 70–90 nt hairpin-shaped precursor (pre-miRNA) by Drosha, a member of the ribonuclease III (RNase III) protein family, and its cofactor, DGC8R, in the context of the microprocessor complex [6]. The pre-miRNA is recognized by the exportin-5-RanGTP heterodimer, and the pre-miRNA is exported to the cytoplasm [18, 33]. In the cytoplasm, pre-miRNA is cut by the RNase III Dicer to generate a ~22 nt miRNA duplex [12, 14]. One strand of the Dicer product remains as a mature miRNA and is assembled into an effector complex known as the miRNA-induced silencing complex (miRISC), which has a gene-silencing effect [15, 26].

RNAi has been used for the functional knockdown of specific proteins in several experimental systems, from cultured cells [7] to whole organisms, including mammals [25]. The RNAi technology represents an interesting tool in reverse genetic studies for several reasons. First, RNAi usually results in partial inhibition of gene expression and thus allows the generation of hypomorphic phenotypes, which are usually impossible to generate by complete knockout via gene targeting [9] or gene editing technologies [31]. Second, the expression of miRNA can be controlled using a large repertoire of RNA pol II-dependent promoters [35], including temporally and spatially specific and inducible promoters [20]. Consequently, the RNA pol II-mediated RNAi strategy can be a powerful alternative to conventional gene knockout technology.

miR-187, an miRNA that is highly expressed in neurons, has been intensively studied, and is considered a potential biomarker in the early diagnosis of a wide range of human cancers [8]. We chose to engineer mir-187 because DraIII and PshAI recognition sites in its precursor sequence allow for the easy replacement of its siRNA.

To confirm the knockdown efficacy of an endogenous gene in vivo, we selected the mouse melanocortin-4 receptor (mMc4r) gene as the target. The mMC4R is a seven-transmembrane G protein-coupled receptor that is predominantly expressed in the hypothalamic nuclei, and is implicated in the regulation of food intake and body weight [2]. Homozygous mMc4r knockout mice are known to express an obese phenotype caused by excess food intake. On the other hand, heterozygous mMc4r knockout mice express an intermediate phenotype, indicating that obesity is tightly coupled to gene dosage [11]. Thus the reduction of mMc4r expression in knockdown mice can be conveniently followed by monitoring body weight.

To test the utility of an mir-187-based shRNA construct driven by an RNA pol II promoter, we used a cytomegalovirus (CMV) early enhancer/chicken β-actin promoter (CAG)-driven expression vector with the potential for expression in a range of tissues and cells [23].

**Materials and Methods**

**Plasmids**

An siRNA-expressing construct was generated as follows. The human miR-187 expression region, containing the mir-187 stem-loop region and 5′ and 3′ flanking regions (271 bp in total), was PCR-amplified from HeLa genomic DNA using the primers 5′-CGCGGATCCATC-GGGGATGCACAGCAAGT-3′ and 5′-GCTC-TAGACCCACCAGAGCCTGGACTTTC-3′, digested with Bsal/HindIII and ligated into the same restriction sites in the pCAG expression vector (pCAG-miR-187). Next, an EF1-α/EGFP expression cassette and an SV40/ blasticidin expression cassette were introduced to visualize the efficacy of transfection and to establish stable cell lines (pGKD-miR-187). Hairpin sequences specific to the targeted mRNAs were inserted into the DraIII/PshAI site of either the pCAG-miR-187 or pGKD-miR-187 vector (pCAG-miR-mMc4r, pCAG-miR-Luc, and pGKD-miR-mMc4r). The inserted oligonucleotide sequences were as follows: mMc4R 5′-GTGACCCCTCCAGGATGCTATTAGCACTTTTTGTGAAGCCACAGATGGAAAAAGTGTGCTCATAAGCCTCGAGGGC-
GC-3′, luciferase: 5′-GTGACCCCTCCAGATTTTCGAGTC
GTCTTAATGTGTGAGCCACAGATGCAATATTAGACGA
AGACGACTCGAAAATCTGAGGGACGC-3′. Underlines indicate modified bases in the 5′ flanking and loop regions. The construct pcDNA3.1-mMC4R contains the mMC4R coding region isolated by PCR.

**Cell culture and transfection**

Human embryonic kidney-derived 293 (HEK293) cells and HeLa cells were grown in DMEM (Thermo Scientific; Logan, UT) supplemented with 10% fetal bovine serum (FBS). Transfections were performed with Lipofectamine Plus (Invitrogen; Carlsbad, CA) as directed by the manufacturer. Cells were co-transfected with pcDNA3.1-mMC4R vector and pCAG vectors at a ratio of 5:1 (wt/wt).

**Mice and ethics statement**

We purchased C57BL/6J mice from the Charles River Laboratory (Yokohama, Japan). Mice were housed under pathogen-free conditions. All procedures involving mice were performed in compliance with National Institutes of Health guidelines and were approved by the Animal Care and Use Committee of Mitsubishi Kagaku Institute of Life Sciences, MITILS. Our animal husbandry abided by MITILS guidelines.

**Generation and genotyping of miR-mMC4R transgenic mice**

Transgenic mice were generated by microinjection into C57BL/6J embryos of the pGKD-miR-mMC4R vector linearized by SalI digestion, using standard techniques [10], and a transgenic line was maintained by crosses with C57BL/6J mice. Transgenic mice were genotyped by Southern blot analysis of tail genomic DNA. In Southern blots, genomic DNAs were digested with NcoI, and a 32P-labelled EGFP fragment was used as a probe (Fig. 1C). We used male mice in whole experiments in this study except the initial screening by weight determination of founder transgenic mice. In addition, we used non-transgenic male littermates as control animals.

**RNA extraction and qPCR**

Total RNA and low-molecular-weight RNA-enriched fractions were isolated using TRIzol reagent (Invitrogen) or an mirVana miRNA isolation kit (Ambion; Austin, TX), following the manufacturers’ protocols.
cDNAs were produced using SuperscriptII Reverse Transcriptase (Invitrogen). The relative expression of mMc4r was estimated by qPCR using SYBR Premix Ex TaqII reagents (TaKaRa; Kyoto, Japan) under standard reaction conditions. The primer sequences were as follows: mouse MC4R-F, 5′-CGCCAGGGTACCAACATGAAG-3′; mouse MC4R-R, 5′-ACATGAAGCACACGAGTATGG-3′; mouse β-actin-F, 5′-GGCCAACCGTGGAAAAGATGA-3′; mouse β-actin-R, CACAGCCTGAGTGCTACGTA-3′. Primers for human GAPDH were purchased from TaKaRa.

Northern blotting for small RNA

In brief, 15% polyacrylamide-8M urea gels were used to isolate the low-molecular-weight RNA-enriched fraction. Band sizes were estimated by comparing them to a 10 bp DNA Marker (NEB; Ipswich, MA). Gels were transferred to Hybond-N+ membranes (GE Healthcare; Uppsala, Sweden) by capillary transfer. After cross-linking, membranes were placed in DIG-easy Hyb (Roche; Indianapolis, IN). DIG-labeled riboprobes were synthesized using an mirVana mirna probe construction kit (Ambion) using the following DNA oligonucleotide as a template: mir-mMC4r sense probe (for guide strand siRNA detection), 5′-AAAAAGTTGCTCATAGCATCCCCTGTCTC-3′. After hybridization at room temperature overnight, membranes were washed twice at 37°C for 15 min in 0.1 × SSC, 0.1% SDS. Probe detection was performed using a DIG Luminescent Detection Kit (Roche) according to the manufacturer’s protocol.

Stripping of the riboprobe was performed at 68°C for 30 min in 0.1% SDS, followed by washing in 2× SSC at room temperature. Then membranes were reprobed with miR-16 riboprobe (Ambion) in the same manner as described above.

Measurement of body weight and food consumption

Body weight was regularly measured, beginning at 3–4 weeks of age. Food intake was measured for male transgenic and non-transgenic mice at 37 to 41 weeks of age. A sufficient amount of food for a 9-day period was weighed and provided to the mice ad libitum. Each weekday morning, the remaining food was weighed, for a total of six measurements.

Determination of metabolic parameters

Fasting male transgenic and non-transgenic mice at 41 weeks of age were sacrificed under anesthesia, and blood, liver, pancreas, and mesenteric fat were collected and weighed. Plasma glucose concentrations were determined using a Glucose-CII Test Kit (Wako Pure Chemical Industries, Osaka, Japan). Insulin concentrations were measured with an enzyme-linked immunosorbent assay (Morinaga Institute of Biologic Science, Yokohama, Japan). Triglyceride levels were measured using a Triglyceride E-test Kit (Wako). Plasma free fatty acid levels were measured using an NEFA C-test Kit (Wako). Plasma leptin levels were measured using Mouse Leptin ELISA kits (Morinaga).

Statistical analysis

Statistical analysis was performed using unpaired two-tailed Student’s t-tests.

Results

Assembly of an siRNA-expression construct based on mir-187 precursor sequences

We constructed an siRNA-expressing cassette containing mir-187 precursor sequences with mir-30 loop sequences (Fig. 1A). Minor modifications to the flanking and loop sequences were introduced to obtain the highest knockdown efficacy by using Dual-Luciferase Reporter Assay system (Fig. 2). In the original shLuciferase cassette with no modification, the G:C pair at the neck of the mir-30 loop was changed to UG wobble pair to open easily in modified siRNA-expressing cassette, Modification (Mod) 1, and 4. As shown in our siRNA design algorithm (22, 29), the 5′ terminal residue of functional siRNA guide strand is preferable to be A or U, and Dicer cleavage site is considered to be at two nucleotides inside from the double strand terminal. Thus, the relative luciferase activity is evidently reduced in Mod.1 cassette compared to the original one, probably because the guide strand is correctly cleaved at A or U at two nucleotides inside from the double strand terminal by this nucleotide modification. Furthermore, AG residues at the two nucleotides downstream of DraIII site in the original mir-187 sequence were changed to C residue in Mod.2, 3, and 4. This region is the binding site of DGCR8, a partner of Drosha, which is a double-stranded RNA binding protein. The accessibility of DGCR8 may increase, since the double stranded region is enlarged by this modification. Indeed, the luciferase activity was also reduced compared to the original one. Then, the expression cassette was inserted into the pCAG
expression vector (pCAG-miR-187, Fig. 1B). Subsequently, an EF1-α/EGFP expression cassette was inserted to visualize the efficiency of transfection in vitro. Furthermore, an SV40/blasticidin gene was inserted for the establishment of stable cell lines. Consequently, the RNAi vector could be utilized for both in vitro and in vivo studies (pGKD-miR-187, Fig.1C). The shRNA sequences against mMc4r or control firefly luciferase (Luc) genes were designed using an algorithm reported previously [22, 29]. The annealed oligonucleotides containing the miR-30 loop sequence were inserted into the pCag-mir-187 or pGKD-mir-187 vector in the site created by a double digest with PshAI and DraIII (pCAG-mir-mMC4R, pCAG-mir-Luc, and pGKD-mir-mMC4R).

Knockdown efficacy of pCAG-miR-mMC4R in cultured cells

We estimated the knockdown efficacy of the pCAG-miR-mMC4R construct against transiently expressed mMc4r in HEK293 cells. The pcDNA3.1/mMC4R expression construct was transiently co-transfected into HEK293 cells with an empty pCag vector without the mir-187 cassette, the pCag-mir-mMC4R construct, or the pCAG-miR-Luc construct. One day after transfection, qPCR analyses were performed to measure mMc4r and human β-actin expression. Co-transfection of the pCAG-mir-mMC4R construct reduced the expression of mMc4r mRNA by 80%, while co-transfection with pCAG-miR-Luc had no effect (Fig. 3). These results indicate that the pCAG-mir-187 expression cassette effectively induced RNAi and specifically repressed target gene expression in vitro.

Generation of GKD-miR-mMC4R transgenic mice

We generated four transgenic founder mice carrying the GKD-miR-mMC4R gene (Suppl. Fig. 1A), and investigated whether the miR-mMC4R construct could knock down endogenous expression of mMc4r in vivo.
Of these transgenic mice, weight gain was observed in only one transgenic male (10L, Suppl. Fig. 1B). Correspondingly, the highest expression among the founder mice of the miR-mMc4r precursor RNA was detected in the hypothalamus of mouse 10L (Suppl. Fig. 2). From these results, we concluded that the obese phenotype observed in 10L can be explained by the greater expression of the miR-mMc4r precursor RNA in the hypothalamus (Suppl. Fig. 2). The body weight of the transgenic mouse line (10L) was further compared to those of wild-type sibs from 6 to 31 weeks of age. A significant increase in the body weight of the transgenic mice started at 11 weeks, and transgenic mice were 30% heavier than non-transgenic littermates at 31 weeks (Fig. 4).

**RNA expression in GKD-miR-mMC4R transgenic mice**

Small RNAs were purified from the hypothalamus, striatum, hippocampus, and pre-frontal cortex regions of the brains of both the GKD-miR-mMC4R transgenic mice (10L) and wild-type sibs, and were examined by Northern blot analysis. Although a faint background signal was also detected in the brain of wild type mouse, a sense probe detected the strong signals of the guide strand of mMc4r siRNA in all of the brain regions extracted from the transgenic mice (Fig. 5). In addition, the mMc4r transcripts in these four brain regions of the transgenic mice were compared to those in the wild-type sibs using qPCR. A significant reduction in the expression of mMc4r mRNA (20–30%) was obtained in all four brain regions of the transgenic mouse line (Table 1). This partial decrement of mMc4r mRNA in the brains of the transgenic mice strongly indicated that the siRNA produced from the GKD-miR-mMC4R transgene attenuated the expression of mMc4r mRNA *in vivo*. 

![Fig. 3. Knockdown efficacy of pCAG-miR-mMC4R in HEK293 cells. HEK293 cells were co-transfected with pcDNA3.1-mMC4R and either pCAG-mir-187 empty vector, pCAG-miR-mMC4R, or pCAG-miR-Luc. One day after transfection, total RNA was extracted and qPCR analyses for mouse Mc4r and human GAPDH were performed. The mean of three independent assays were shown. Each error bar represents the SE.](image)

![Fig. 4. Weight gain of GKD-miR-mMC4R transgenic mice. The symbols indicate the first week that body weight differed significantly between transgenic (Tg; n=9) and non-transgenic (nTg; n=15) mice (*; P<0.05, †; P<0.01). Error bars indicate SE.](image)
Food consumption and metabolic phenotypes of GKD-miR-mMC4R transgenic mice

Next, we examined the hyperphagic phenotype in male litters of transgenic mice. The total food consumption of 37- to 41-week-old transgenic mice was determined over 9 days. The knockdown of mMc4r resulted in a significant increase (9%) in normal diet consumption over that of non-transgenic littermates (Table 2). To further understand the obese phenotype of transgenic mice, we performed a number of assays of parameters related to obesity: oral glucose tolerance tests (OGTT); measurements of glucose, triglyceride (TG), insulin, free fatty acid (FFA), total cholesterol, and leptin in serum; measurement of tissue weight for mesenteric fat, pancreas, and liver; and measurement of leptin and TG levels in the latter two tissues, respectively. There were no significant differences in glucose, TG, FFA, or total cholesterol levels between fasting transgenic and non-transgenic mice. However, insulin and leptin were significantly elevated in transgenic mice, by 6-fold and 2.5-fold, respectively.

In transgenic mice, the tissue weights of pancreas, liver, and mesenteric fat were significantly increased. The content of insulin in pancreas showed a 3.5-fold increase in transgenic mice. Furthermore, liver TG levels were also significantly elevated in transgenic mice (Table 2). The AUC0-2h of the glucose response during OGTT tended to increase in transgenic mice compared to non-transgenic mice but was not significantly different (Suppl. Fig. 3).

From these observations, we concluded that GKD-miR-mMC4R transgenic mice had a phenotype of hyperinsulinemia, fatty liver, functional disorder of the pancreas, and the accumulation of mesenteric fat.

In conclusion, we have determined that a mir-187-based expression construct is practically useful for both in vitro and in vivo RNAi. The expression cassette introduced into mice could express functional siRNAs against target genes in tissues. While transgenic miR-mMC4R mice had the obese phenotype previously seen in knockout mice, this has revealed that such a phenotype may be the result of as little as a 20–30% partial knockdown of mMc4r expression in vivo.

**Discussion**

We have described the development of a new siRNA-expressing cassette, and applied it in RNAi in vitro and in vivo. The method depends on the expression of specific siRNAs within mir-187 precursor sequences. Because mir-187 has recognition sites for the restriction enzymes DraIII and PshAI in its flanking regions, syn-
thetic hairpin DNA sequences can be directly inserted into the vectors (Fig. 1A). This enables high-throughput construction of siRNA-expressing libraries.

A previous study showed that the body weights of homozygous mC4r mutant mice were 50% heavier than those of wild-type mice at 15 weeks of age, and heterozygous mice showed an intermediate weight gain of approximately 25% [11]. In the present study, miR-mMC4r transgenic mice, whose mRNA level of mC4r decreased by 20~30% (Table 1), body weights were 15% heavier than those of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Additionally, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age (Fig. 4). Furthermore, in the abovementioned previous study, food consumption of homozygous mC4r knockout mice increased by 40% over that of wild-type littermates at 15 weeks of age.

Table 2. Food consumption and metabolic phenotypes of GKD-miR-mMC4R transgenic mice

|                          | non-Tg (n=9) | Tg (n=15) |
|--------------------------|-------------|-----------|
| **Food consumption**     |             |           |
| Total food intake for 9 days (g) | 35.6 ± 3.4 | 38.4 ± 3.3 |
| **Serum**               |             |           |
| Glucose (mg/dl)          | 85.12 ± 3.20| 89.12 ± 3.76|
| Triglyceride (mg/dl)     | 71.8 ± 1.8  | 73.6 ± 3.6 |
| Insulin (ng/ml)          | 3.2 ± 0.3   | 20.0 ± 6.7 |
| Free fatty acid (uEq/l)  | 1,369.3 ± 44.4| 1,483.4 ± 98.0|
| Total cholesterol (mg/dl)| 181.9 ± 10.8| 199.4 ± 11.6|
| Leptin (ng/ml)           | 19.8 ± 2.9  | 50.8 ± 15.7 |
| **Pancreas**            |             |           |
| Insulin (ng/ml/g tissue) | 58,387.2 ± 3,246.2| 203,192.5 ± 28,998.7|
| Tissue weight (g)       | 0.372 ± 0.013| 0.399 ± 0.016|
| **Liver**               |             |           |
| Triglyceride (mg/g tissue) | 28.3 ± 1.3  | 38.1 ± 2.0 |
| Tissue weight (g)       | 1.387 ± 0.060| 2.258 ± 0.174|
| **Mesenteric Fat**      |             |           |
| Tissue weight (g)       | 0.990 ± 0.067| 1.653 ± 0.070|

Mean ± SE, *; P<0.05, **; P<0.01 (t-test).

Although the miR-mMC4R expression construct mediates effective (around 80%) knockdown in cell culture (Fig. 3), the knockdown efficiency in transgenic mice was limited to about 20~30% (Table 1). One potential reason for this discrepancy is that the expression level of transgenic siRNA against mC4r was not enough to achieve effective knockdown in vivo. In Northern blot analysis, approximately 1 ng siRNA was detected in 6 μg small RNA extracted from striatum (Fig. 5). The 6 μg small RNA was obtained from about 80 mg striatal tissue (data not shown). The concentration of 1 ng siRNA / 80 mg tissue is comparable with ~1 nM siRNA. There are two main reasons that explain the low-level expression of transgenic siRNA. One possibility is that a large amount of shRNA expressed from the Cag promoter could saturate the activity of exportin-5, a rate-limiting factor required for nuclear export of pre-miRNAs [5]. Such high shRNA expression may inhibit endogenous miRNA functions required for normal embryogenesis and development [30]. Consequently, only transgenic mice with low shRNA expression would survive to birth. In fact, after microinjection of the Cag-EGFP transgene, bright EGFP expression in embryos can be observed at early developmental stages, from the 2-cell stage to blastocysts [13]. To overcome this possible problem, conditional expression constructs, with inducible or stage-specific RNA pol II promoters, may be valuable.

Another possible reason for the low-level expression of transgenic siRNA is that the silencing of transgene by the vector sequences or by the random and multi-copy integration [4, 24, 28]. Especially, the vector used in this study (Fig. 1C) contained prokaryote-derived elements from pBluescript, therefore, it’s plausible to remove such bacterial sequences before injection into mice eggs. Although the obese phenotypes of miR-mMC4R transgenic mice were similar to those of mC4r knockout mice in terms of hyperinsulinemia [1] and fatty liver...
[27], the serum levels of glucose were normal in miR-
mMC4R transgenic mice (Table 2). In previous studies, pre-obese mMc4r homozygous knockout mice did not display hyperglycemia and hyperinsulinemia, although they were already significantly heavier than heterozy-
gous knockout and wild-type mice [1]. mMc4r knockout mice develop both hyperglycemia and hyperinsulinemia [11], whereas the miR-mMC4R transgenic mice in our study exhibited only hyperinsulinemia (Table 2). Re-
expression of mMc4r specifically in the lateral hypotha-
lamic area (LHA) improves glucose metabolism in obese 
mMc4r knockout mice, without affecting body weight or circulating insulin levels [21]. It is possible that a partial knockdown of mMc4r (20% to 30%) did not affect LHA signaling in the regulation of glucose homeostasis. The partial knockdown by the mir-187 expression con-
struct in vivo may be a powerful tool to investigate the relationship between the level of gene expression and the phenotype of interest.

In conclusion, we developed a new siRNA-expression construct based on the mir-187 precursor. Transgenic mice with partial knockdowns offer valuable functional information. Such mir-187 precursor-based constructs will provide new tools for validating drug target genes in vitro and in vivo, and for generating hypomorphic animal models.

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