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ORIGINAL ARTICLE

MicroRNA-21 is a potential link between non-alcoholic fatty liver disease and hepatocellular carcinoma via modulation of the HBP1-p53-Srebp1c pathway

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ABSTRACT

Background Non-alcoholic fatty liver disease (NAFLD) is a major risk factor for hepatocellular carcinoma (HCC). However, the mechanistic pathways that link both disorders are essentially unknown.

Objective Our study was designed to investigate the role of microRNA-21 in the pathogenesis of NAFLD and its potential involvement in HCC.

Methods Wildtype mice maintained on a high fat diet (HFD) received tail vein injections of microRNA-21-antisense oligonucleotide (ASO) or miR-21 mismatched ASO for 4 or 8 weeks. Livers were collected after that time period for lipid content and gene expression analysis. Human hepatoma HepG2 cells incubated with oleate were used to study the role of miR-21 in lipogenesis and analysed with Nile-Red staining. microRNA-21 function in carcinogenesis was determined by soft-agar colony formation, cell cycle analysis and xenograft tumour assay using HepG2 cells.

Results The expression of microRNA-21 was increased in the livers of HFD-treated mice and human HepG2 cells incubated with fatty acid. MicroRNA-21 knockdown in those mice and HepG2 cells impaired lipid accumulation and growth of xenograft tumour. Further studies revealed that *Hbp1* was a novel target of microRNA-21 and a transcriptional activator of p53. It is well established that p53 is a tumour suppressor and an inhibitor of lipogenesis by inhibiting Srebp1c. As expected, microRNA-21 knockdown led to increased HBP1 and *p53* and subsequently reduced lipogenesis and delayed G1/S transition, and the additional treatment of HBP1siRNA antagonised the effect of microRNA-21-ASO, suggesting that HBP1 mediated the inhibitory effects of microRNA-21-ASO on both hepatic lipid accumulation and hepatocarcinogenesis. Mechanistically, microRNA-21 knockdown induced p53 transcription, which subsequently reduced expression of genes controlling lipogenesis and cell cycle transition. In contrast, the opposite result was observed with overexpression of microRNA-21, which prevented *p53* transcription. **Conclusions** Our findings reveal a novel mechanism by which microRNA-21, in part, promotes hepatic lipid accumulation and cancer progression by interacting with the Hbp1-p53-Srebp1c pathway and suggest the potential therapeutic value of microRNA-21-ASO for both disorders.

INTRODUCTION

The incidence of hepatocellular carcinoma (HCC) worldwide nearly matched its mortality,

Significance of this study

What is already known on this subject?

- miR-21 is upregulated in human hepatocellular carcinoma.
- ▶ p53 is a transcriptional repressor of *Srebp1c*.

What are the new findings?

- miR-21 is highly expressed in hepatocytes, and its expression is significantly increased in livers of dietary obese mice and human HepG2 cells incubated with fatty acid.
- Antagonising miR-21 in liver prevents hepatic lipid accumulation and growth of xenograft tumour.
- miR-21 knockdown prevents G1/S transition and cancer cell proliferation.
- ► *HBP1* is a novel target of miR-21 and a transcriptional activator of *p53*.
- HBP1 mediates the inhibitory effects of miR-21-anti-sense oligonucleotide on hepatic lipid accumulation and hepatocarcinogenesis.
- miR-21 is a potential association between non-alcoholic fatty liver disease (NAFLD) and hepatocellular carcinoma (HCC) via interacting with the *Hbp1-p53-Srebp1c* pathway.

How might it impact on clinical practice in the foreseeable future?

 Our data suggest that miR-21 is a potential therapeutic target for both NAFLD and HCC.

demonstrating the aggressive nature of this malignancy and limited therapeutic options.¹ Although HBV and HCV are major risk factors of HCC, nonalcoholic fatty liver disease (NAFLD) remains a common underlying pathology to the majority of patients with HCC in the Western world.² The incidence of NAFLD is growing rapidly due to the prevalence of obesity.³ It is estimated that 90% of obese patients have some form of fatty liver, ranging from simple steatosis to more severe forms of non-alcoholic steatohepatitis (NASH) and cirrhosis with its associated high risk of HCC. In addition, given limited effects of chemotherapy and the relative insensitivity of HCC to radiotherapy, complete tumour extirpation represents the only choice for a long-term cure. Unfortunately, the



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majority of patients are not eligible for surgical resection because of tumour extent or underlying liver dysfunctions including NAFLD. As described above, despite the strong association between NAFLD and HCC, the underlying mechanisms are largely unknown due in part to their complex nature of disease.

The discovery of a class of naturally occurring small noncoding RNAs, termed microRNAs (miRNAs),^{4 5} has stimulated a new field of research on NAFLD and HCC. Alterations in miRNA expression have been reported in human individuals with NAFLD/NASH and HCC.⁶ ⁷ Reflective of their key roles in lipid metabolism and carcinogenesis,⁵ ⁸ miRNAs have been suggested as novel therapeutic targets for both metabolic diseases and human cancers. However, the miRNAs associated with both NAFLD and its potential sequel HCC are poorly described. Our interest in miR-21 arose initially from hepatocyte-specific miRNA profiling studies in mouse livers, in which we showed that miR-21 is highly expressed in hepatocytes. Furthermore, we observed that high fat diet (HFD) treatment significantly induced expression of miR-21 in livers of mice. By antagonising miR-21 in liver, we were able to prevent hepatic lipid accumulation in dietary obese mice. Consistent with our findings, miR-21 expression was significantly upregulated in human patients with NASH.⁶ It is also known that miR-21 is a potent promoter of HCC and other human cancers.^{7 9} These data led us to hypothesise that miR-21 plays an important role in the pathogenesis of NAFLD and its potential progression to HCC. In the present study, we have investigated the regulatory role of miR-21 in linking NAFLD and HCC in both in vivo and in vitro model systems.

MATERIALS AND METHODS

Bioinformatic analysis

Identification of miR-21 target genes was conducted as previously described with minor revision.¹⁰ In detail, we compiled a list of downregulated genes in livers of patients with NAFLD/ NASH by downloading their microarray data from GEO (http:// www.ncbi.nlm.nih.gov/geo/).¹¹ mRNA profiles of six normal liver samples (male) and eight NAFLD/NASH liver samples (male) were compared using GeneSpring (Agilent Technologies Genomics). Differentially expressed genes were defined by a log-scale ratio ≤ 0.3 between paired samples with a p< 0.05. Based on these criteria, we identified 1219 downregulated probes in NAFLD/NASH samples (see online supplementary table S1). To identify genes with binding motifs for miR-21, we downloaded the target gene databases of miR-21 based on TargetScan,¹² Pictar¹³ and Starbase.¹⁴ Only hits from Target or PicTar algorithm that were confirmed by Ago HITS-CLIP (highthroughput sequencing of RNAs isolated by cross-linking immunoprecipitation (HITS-CLIP) from Argonaute protein complex) were selected. These three databases were compared using Microsoft Access 2000, yielding 219 potential targets that have miR-21 binding motif (see online supplementary table S2). We then compared 1219 downregulated probes in livers of patients with NAFLD/NASH with 219 genes that have at least one binding motif for miR-21 using Microsoft Access 2010, which resulted in an overlap of 13 genes between two databases that were considered as potential targets of miR-21 (see online supplementary table S3). Gene ontology (GO) analysis was done using PathwayStudio software (Elsevier).

Animal, diet treatment and sample collection

Male $Dicer1^{fl/fl}$ mice on a mixed 129S4, C57Bl/6 strain background¹⁵ were crossed with C57Bl/6 *Alb-Cre*^{+/-} mice¹⁶ to generate $Dicer1^{fl/fl}$, *Alb-Cre*^{+/-} mice (mice are from Dr Holger Willenbring's lab at the University of California, San Francisco). To specifically investigate the impact of miRNAs on mature liver function, we initiated Cre recombinase expression in 8-week-old to 10-week-old mice.¹⁷ To restrict *Cre* expression to hepatocytes, we used a hepatocyte-specific *Transthyretin* (*Ttr*) promoter and pseudotyped the vector genome with capsids from AAV8, a serotype that can transduce virtually all hepatocytes in vivo without causing toxicity.^{10 17 18}

To determine the effect of hepatic lipid accumulation on miRNA expression, 8-week-old wildtype male C57Bl/6 mice (Jackson Laboratory, n=6) were maintained on either a normal chow diet (Open Source D12450B: 10% Kcal fat) or an HFD (Open Source D12492: 60% Kcal fat) for 4 weeks as described by Vickers *et al.*¹⁹ After 4 weeks of HFD administration, livers were collected for miRNA and gene expression analysis.

To determine the role of miR-21 in NAFLD, we synthesised locked nucleic acid anti-miR-21 anti-sense oligonucleotide (ASO) (Exiqon) specifically targeting miR-21 and also generated miR-21-mismatched-ASO (miR-21-MM-ASO), a control ASO that differs from miRNAs in four mismatched base pairs. The male C57Bl/6 mice kept on HFD for 4 weeks were divided into two groups; one group (n=8) was treated with miR-21-ASO and the other with miR-21-MM-ASO (control, n=8). Mice received a dose of 25 mg/kg miR-21-ASO or miR-21-MM-ASO (0.9% NaCl) weekly for 4 or 8 weeks via tail vein injection. At those times, the mice were anaesthetised, and blood was collected by way of cardiac puncture. Subsequently, the livers were harvested and immediately frozen in liquid nitrogen and stored at -80° C for gene expression and histological analysis.

Fatty acid treatment of HepG2 cells

HepG2 cells were obtained from Dr Xin Chen's laboratory at the University of California, San Francisco. Sodium oleate was obtained from Sigma-Aldrich and was dissolved in Dulbecco's modified Eagle medium (DMEM) with 1% fatty acid free bovine serum albumin (BSA) (Sigma). Oleate treatment of HepG2 cells was carried out as previously described with minor revision.^{10 20} Specifically, HepG2 cells were plated in four-well chamber slides with DMEM medium supplemented with 10% fetal bovine serum (Invitrogen). After 24 h, HepG2 cells were treated with either control medium (DMEM supplemented with 1% fatty acid free BSA) or medium containing oleate (0.5 mM). The cells were cultured for another 24 h, then lipid accumulation and miR-21 expression were determined by Nile Red Staining (Sigma-Aldrich) and qRT-PCR, respectively (see online supplementary materials and methods for details).

Cell proliferation analysis

HepG2 cells were transfected with miR-21-ASO, scrambled control or miR-21-ASO plus *HBP1*-siRNA using lipofectamine 2000 (Invitrogen). After 48 h, cell proliferation was determined using a MTT (3-(4,5-dimethylthiazol-2-yl)-2, 5-diphenyltetrazoliumbromide) cell proliferation kit (Cell Biolabs) according to the manufacturer's instruction (see online supplementary material and methods for details).

Focus formation assay and flow cytometry analysis

HepG2 cells were used to determine the effect of miR-21 on focus formation and cell cycle progression (see online supplementary material and methods for details).

Xenograft tumour assay

Male BALB/c athymic nude mice (6 weeks old) purchased from Jackson Laboratory were used to study the role of miR-21 in

promoting growth of xenograft tumour from HepG2 cells (see online supplementary material and methods for details).

Histological analysis

Frozen sections of liver were stained with Oil Red-O staining. Paraformaldehyde-fixed, paraffin-embedded sections of liver were stained with H&E (see online supplementary material and methods for details).

Lipid and lipoprotein analysis

Both plasma and hepatic lipid content were enzymatically measured in liver lysates and plasma via a colorimetric assay using a triglyceride assay kit from Roche Diagnostics, according to the manufacturer's protocols (see online supplementary material and methods for details).

Western blot and Q-RT-PCR

Western blot and qRT-PCR were used to determine expression levels of genes. Primers used for quantitative RT-PCR are listed in online supplementary table S4 (see online supplementary material and methods for details).

Statistical analysis

Statistical analysis was performed using GraphPad Prism Software. Data derived from cell-line experiments were presented as mean \pm SD and assessed by a two-tailed Student's t test. Statistical difference for cell cycle progression analysis was evaluated using χ^2 test. Mann–Whitney test was used to evaluate the statistical significance for mouse experiments. Each experiment was repeated at least three times; and the error bars represent the SD. p<0.05 was considered to be statistically significant.

RESULTS

miR-21 is robustly induced in livers of mice on HFD and HepG2 cells exposed to high levels of fatty acid

Hepatocytes are the major cells that control lipid metabolism and the primary site of NAFLD and HCC. To investigate the role of miRNAs in both disorders, we compiled hepatocytespecific miRNA profiles by comparing miRNAs expression of livers of hepatocyte-specific *Dicer1* knockout (*DKO*) and wildtype mice (see online supplementary table S5). We observed that miR-21 was the most significantly downregulated miRNA in livers of *DKO* mice (\geq 39-fold reduced), indicating that hepatocytes represent a main source of its expression in the liver (figure 1A). To assess the role of miR-21 in NAFLD, we fed wildtype C57Bl/6 mice an HFD (see online supplementary figure S1A–C) and measured its hepatic expression. The results showed that miR-21 had a twofold upregulation in the livers of HFD-treated mice (figure 1B), suggesting its potential role in NAFLD.

We also determined whether fatty acids can increase the expression of miR-21 in human hepatoma HepG2 cells. Oleic acids are the most abundant unsaturated fatty acids in liver triglycerides in both normal subjects and patients with NAFLD.²¹ In this study, HepG2 cells were used because of their increased sensitivity to fat accumulation.²¹ Nile-Red staining revealed that oleic acid treatment led to an increase in intracellular lipids in HepG2 cells (figure 1C, D), which was also associated with upregulation of miR-21 (figure 1E). Taken together, our in vivo and in vitro studies indicated that both HFD treatment of mice and exposure of HepG2 cells to fatty acid were able to induce expression of miR-21.



Figure 1 Hepatocyte-specific miR-21 is significantly induced in livers of high fat diet (HFD)-treated mice and HepG2 cells treated with fatty acid. (A) gRT-PCR confirmed that miR-21 was highly expressed in hepatocytes of liver. miR-21 expression in liver was compared between Dicer1 knockout (DKO) (n=3) and wildtype mice (n=3) using qRT-PCR. Data are presented as mean±SD (p=0.0016, Student's t test). (B) HFD treatment led to higher levels of miR-21 in livers of mice. Briefly, wildtype mice (C57Bl/6) were maintained on HFD for 4 weeks, and then mice were sacrificed and livers collected for miRNA expression analysis. Control mice received standard diet (SD). Data are presented as mean ±SD (p=0.0022, Mann-Whitney test). (C and D) Oleate treatment increased lipid content and subsequently (E) expression of miR-21 in HepG2 cells. HepG2 cells were maintained in Dulbecco's modified Eagle medium containing 0.5 mM oleate. Data are presented as mean±SD. In this multiple-groups experiment, we only performed comparison between two groups and Student's t test was used for statistical analysis. Lipid droplets in human hepatocytes were labelled with arrows.

To further elucidate the role of miR-21 in hepatic lipid accumulation, we began to identify target genes of miR-21 by combining mRNA profiling of livers of NAFLD individuals with the bioinformatic prediction of miR-21 binding motifs within potential target mRNAs. This led us to identify 13 genes including *HBP1*, *SOX7* and *RHOB* that showed reduced expression in human fatty liver and contained a conserved binding motif for miR-21 (see online supplementary table S3). GO analysis of the above 13 genes revealed that *HBP1* was a potent tumour suppressor by preventing G1/S transition of cell cycle.^{22 23} In addition, our prediction from in silico algorithms showed that the 3' UTR of *HBP1* mRNA was 100% complementary to the miR-21 *5'* seed region, exhibiting the highest prediction score and binding energy (figure 2A). Therefore, we selected *HBP1* as a potential target of miR-21.



Figure 2 Hbp1 is a direct target of miR-21. (A) Bioinformatic analysis showing that the seed sequence of miR-21 has a high level of complementarity to Hbp1 3' UTR, prediction score and favourable binding energy. Complementary sequences to the seed region of miR-21 within the 3' UTRs of Hbp1 are conserved between human, mouse and monkey (grey highlight). (B) miR-21 knockdown in high fat diet (HFD)-treated mice led to increased Hbp1 mRNA levels in liver. C57BI/6 wildtype mice were kept on normal chow until 8 weeks of age and then maintained on HFD until 16 weeks of age. At 12 weeks of age, the mice were given miR-21-anti-sense oligonucleotide (ASO) (25 mg/kg, tail vein injection) until 16 weeks of age. C57BI/6 mice maintained on HFD and treated with miR-21-MM-ASO served as controls. The expression levels of Hbp1 were determined by qRT-PCR. Data are presented as means±SD (p=0.0002, Mann-Whitney test). (C) miR-21 mimic transfection into Hepa1,6 cells caused dose-dependent inhibition of the activity of a luciferase reporter gene linked to the 3' UTR of mouse Hbp1. Data are presented as mean±SD. In this multiple-groups experiment, we only performed comparison between two groups among them and Student's t test was used for statistical analysis. (D) Conversely, transfection with a miR-21 inhibitor antagonised the binding of miR-21 mimics to the 3' UTR of mouse Hbp1, which was reflected by increased luciferase activity. Data are presented as mean±SD. Student's t test was used for statistical analysis. (E) Mutated binding motif for miR-21 within Hbp1 3'UTR impaired miR-21 binding, which was reflected by (i) negligible change of luciferase activity after miR-21 mimics treatment; and (ii) miR-21 inhibitor treatment had no effect on luciferase activity. Data are presented as mean \pm SD (p \geq 0.1, Student's t test). (F) Transfection of miR-21 mimics into HepG2 cells inhibited expression levels of endogenous HBP1 as revealed by gRT-PCR and western blot. Data are presented as mean±SD (p=0.0041, Student's t test). (G) miR-21 knockdown by transfecting miR-21 inhibitor into HepG2 cells caused an increase in endogenous HBP1 at the protein and mRNA levels. Data are presented as mean±SD (p=0.0002, Student's t test).

We next determined its expression in the livers of dietary obese mice treated with miR-21-ASO. It was not surprising that Hbp1 expression increased more than twofold in the livers of miR-21-ASO-treated mice compared with those treated with miR-21-MM-ASO (figure 2B). Taken together, hepatic expression of miR-21 was increased in dietary obese mice and livers of human patients with NAFLD/NASH, and Hbp1, as a potential target of miR-21, showed reduced expression in livers of obese mice and human patients with NAFLD (see online supplementary table S3 and figure S1D). Our findings suggested that the crosstalk between miR-21 and Hbp1 might play an important role in hepatic lipid accumulation.

Hbp1 is a direct target of miR-21

To establish that miR-21 directly recognises the predicted target site within the 3' UTR of *Hbp1*, the 3' UTR of mouse *Hbp1* mRNA was cloned into a luciferase reporter vector (pMiR-Report) to generate pMiR-*Hbp1*. Mouse Hepa1,6 cells were transfected with pMiR-*Hbp1* and chemically synthesised miR-21 mimic or miR-21 inhibitor. We found that miR-21 mimics significantly downregulated luciferase activity in a dosedependent fashion (figure 2C). Consistently, miR-21 inhibitor antagonised the inhibitory effect of miR-21 mimics on luciferase activity (figure 2D). Furthermore, we mutated the binding motif for miR-21 within the pMiR-*Hbp1* 3' UTR and found that both mimics and inhibitors of miR-21 had no effect on luciferase activity (figure 2E), indicating a potentially direct interaction between miR-21 and *Hbp1* mRNA. To further validate that *Hbp1* is a target of miR-21, we increased intracellular levels of miR-21 in HepG2 cells in the absence of fatty acid. qRT-PCR and western blot revealed that miR-21 significantly inhibited expression of *HBP1* (figure 2F). In contrast, miR-21 knockdown led to an increase in mRNA and protein levels of *HBP1* in HepG2 cells (figure 2G). Together, these results confirmed that *Hbp1* is a direct target of miR-21.

HBP1 inhibits expression of *SREBP1C*, *CCND1* and *CCNB1* by activating *p53*

HBP1 is a well-described transcriptional repressor that modulates expression of genes involved in cell cycle progression.²³ Therefore, we overexpressed *HBP1* in HepG2 cells and determined the expression levels of genes involved in cell cycle using Human Cell Cycle RT² Profiler PCR Assay. Interestingly, we observed that *p53* was the most upregulated after overexpression of *HBP1* (figure 3A). *p53* functions as a tumour suppressor and potent inhibitor of lipogenesis by inhibiting transcription of *SREBP1C*,^{24–26} leading to our hypothesis that miR-21 plays roles in both lipogenesis and carcinogenesis by interacting with the HBP1-p53 pathway.

Overexpression of HBP1 led to increased mRNA levels of p53, implying that HBP1 might activate transcription of p53 by binding to its promoter. Therefore, we cloned the *p53* promoter into a luciferase reporter vector (pGL3-Basic) and generated pGL3-p53. Hepa1,6 cells were transfected with pGL3-p53 and HBP1 expression vector. As expected, overexpression of HBP1 induced luciferase activity (figure 3B), and HBP1 knockdown led to decreased luciferase activity (figure 3C). Furthermore, HBP1 knockdown impaired expression of endogenous p53 (figure 3D), suggesting that HBP1 was able to activate transcription of p53. HBP1 can function as a transcriptional activator by binding to a specific binding motif (GGGATGGG).²² However, we did not identify this binding motif within the promoter of p53, signifying that HBP1 might activate transcription of p53 by interacting with other transcription factors that have binding sites within the *p53* promoter.

Srebp1c is a transcription factor that activates genes encoding enzymes required for lipid synthesis.^{27 28} Considering the role that p53 plays in inhibiting lipogenesis by modulating Srebp1c,²⁶ we cloned the mouse Srepb1c promoter into pGL3-basic vector (pGL3-Srebp1c). As expected, co-transfection of p53 expression vector pGL3-Srebp1c and into Hepa1,6 cells significantly reduced luciferase activity in a dose-dependent fashion (see online supplementary figure S2A), and in contrast, p53 knockdown induced luciferase activity (see online supplementary figure S2B). Furthermore, we observed that overexpression of p53 led to a decrease in endogenous mRNA levels of SREBP1C and its targeted lipogenic genes SCD1 (stearoyl-CoA desaturase-1), GPAT (glycerol 3-phosphate acyltransferase), and FASN (fatty acid synthase),²⁹ and genes controlling cell cycle progression including CCNB1 and CCND1 in HepG2 cells (figure 3E).³⁰ p53 knockdown led to the opposite effect (figure 3F), underscoring the central role of p53 in modulating the expression of genes involved in lipogenesis and cell cycle progression.

To further determine whether *HBP1* prevents transcription of genes associated with lipogenesis and G1/S transition via *p53-SREBP1C* pathway, we overexpressed *HBP1* in HepG2 cells and determined expression levels of *p53*, *SREBP1C*, the lipogenic genes and *CCNB1* and *CCND1*. As confirmed by qRT-PCR, *HBP1* overexpression led to increased *p53*, which

subsequently prevented expression of *SREBP1C* and *SCD1*, *FASN* and *GPAT*, as well as *CCNB1* and *CCND1* (figure 3G). In summary, our findings suggested that HBP1 is able to simultaneously inhibit expression of *CCNB1*, *CCND1* and *SREBP1C* by modulating *p53*.

miR-21 prevents expression of *p53* but promotes transcription of *SREBP1C* by modulating *HBP1* expression

We have shown that *HBP1* is a target of miR-21, and HBP1 can activate transcription of p53. Meanwhile, p53 is a transcriptional repressor of *Srebp1c*.²⁵ ³¹ Thus, we hypothesised that miR-21 can simultaneously regulate expression of genes involved in lipogenesis and the G1/S transition by modulating the *HBP1-p53-SREBP1C* pathway. Indeed, overexpression of miR-21 in HepG2 cells inhibited expression of *HBP1*, which subsequently led to a reduction in *p53* and an increase in mRNA levels of *CCND1*, *CCNB1* and *SREBP1C*, as well as its target genes including *SCD1*, *FASN* and *GPAT* (figure 3H), while miR-21 knockdown led to an opposite effect (figure 3I). Our findings indicated that miR-21 is able to modulate expression of genes controlling lipogenesis and cell cycle progression via the p53-SREBP1C pathway.

miR-21 modulates lipid accumulation in HepG2 cells by interacting with the *HBP1-p53* pathway

We then determined whether overexpression of miR-21 can promote lipogenesis. As expected, miR-21 overexpression prevented expression of *HBP1* and *p53* (figure 4A), which subsequently promoted lipid accumulation in HepG2 cells (figure 4B, C).

To determine loss of function for miR-21 in lipid accumulation, we transfected miR-21 inhibitor into oleate-treated HepG2 cells to knock down upregulated miR-21. Antagonising miR-21 led to a significant increase in HBP1 and p53 (figure 4D), which subsequently prevented lipid accumulation (figure 4E, F). These data demonstrated that miR-21 was sufficient for the downregulation of HBP1 and p53, which subsequently induced lipid accumulation. To further investigate the role of the interaction between miR-21 and HBP1 in hepatic lipid accumulation, we mutated the binding motif for miR-21 within the 3' UTR of Hbp1 in the pMiR-Hbp1 (referred to as pMiR-Hbp1Mu) and introduced the pMiR-Hbp1 or pMiR-Hbp1Mu into oleatetreated HepG2 cells. Since oleate treatment increases miR-21 expression in HepG2 cells, it was expected that it would lead to a decrease of luciferase activity in HepG2 cells transfected with pMiR-Hbp1 compared with pMiR-Hbp1Mu. In fact, oleate treatment of HepG2 cells transfected with pMiR-Hbp1 resulted in robust repression of luciferase activity compared with pMiR-Hbp1-Mu (figure 4G). Together, our results indicated that HBP1 is a direct target of miR-21 during lipid accumulation in HepG2 cells and the crosstalk of miR-21 with HBP1 and p53 plays an important role in hepatic lipid accumulation.

Inhibitory effect of miR-21-ASO on hepatic lipid accumulation is mediated by *HBP1*

To confirm that miR-21 promotes lipogenesis via *HBP1*, we initially antagonised miR-21 by transfecting miR-21-ASO into HepG2 cells to induce expression of *HBP1*, and then knocked down the induced *HBP1* using *HBP1*-siRNA. The results showed that miR-21 knockdown increased *p53* and *HBP1* and reduced lipid content in HepG2 cells, but additional treatment of *HBP1*-siRNA offset the effect of miR-21-ASO (figure 5A–C), which suggested that *HBP1*, in part, mediated the inhibitory effect of miR-21 inhibitor on lipid accumulation. To study the



miR-21 inhibitor

Figure 3 HBP1 is a transcriptional activator of p53. (A) Overexpression of HBP1 increased mRNA levels of p53 in human HepG2 cells. Data are presented as means±SD (p=0.0003, Student's t test). (B) Overexpression of HBP1 caused dose-dependent increase of the activity of a luciferase reporter gene linked to the p53 promoter. Data are presented as mean±SD. Student's t test was used for statistical analysis. (C) HBP1 knockdown via its siRNA decreased luciferase activity driven by p53 promoter. Data are presented as means±SD. Student's t test was used for statistical analysis. (D) HBP1 knockdown via its siRNA resulted in decreased mRNA levels of p53. Data are presented as mean±SD (*p=0.0040, Student's t test). (E) Overexpression of p53 led to deceased SREBP1C, lipogenic genes SCD1, FASN and GPAT as well as CCND1 and CCNB1 in HepG2 cells. Data are presented as mean \pm SD (*p<0.05; **p<0.001; Student's t test). (F) Knockdown of *p53* via its siRNA increased mRNA levels of *SREBP1C*, *FASN*, *SCD1*, *GPAT*, *CCND1* and *CCNB1* in HepG2 cells. Data are presented as mean \pm SD (*p<0.05; **p<0.001; Student's t test). (G) Overexpression of *HBP1* reduced expression of *p53*, which subsequently led to decreased expression of genes involved in lipogenesis and G1/S transition. Data are presented as mean±SD (*p<0.05; **p<0.001; Student's t test). (H) Overexpression of miR-21 inhibited HBP1 and p53, which subsequently promoted expression of the lipogenic genes including SCD1, FASN and GPAT and the genes controlling cell cycle progression including CCNB1 and CCND1. Data are presented as mean±SD (*p<0.05; **p<0.001; Student's t test). (I) miR-21 knockdown via its inhibitor led to increased HBP1 and p53 and decreased expression of the lipogenic and cell cycle-related genes. Data are presented as mean±SD (*p<0.05; **p<0.001; Student's t test). Figure 4 MiR-21 modulates lipid accumulation in HepG2 cells by interacting with the HBP1-p53 pathway. (A) Overexpression of miR-21 inhibited expression of HBP1 and p53, which (B and C) subsequently promoted lipid accumulation in HepG2 cells in the presence of 0.25 mM oleate. Data are presented as mean ±SD. Student's t test was used for statistical analysis. Lipid droplets were labelled with arrows. (D) miR-21 inhibitor transfection into HepG2 cells cultured with the medium containing 0.5 mM oleate led to an increase in HBP1 and p53, which (E and F) antagonised the effect of upregulated miR-21 on lipid accumulation. Lipid droplets were labelled with arrows. Data are presented as mean±SD. Student's t test was used for statistical analysis. (G) Oleate treatment led to a decrease in luciferase activity of pMiR-Hbp1 as compared with pMiR-Hbp1Mu. Data are presented as mean±SD (p=0.0030, Student's t test).



role of the crosstalk between p53 and miR-21 in lipogenesis, we transfected oleated-incubated HepG2 cells with miR-21 mimics or a combination of miR-21 mimics and p53 expression vector. Nile-Red staining and qRT-PCR revealed that miR-21 mimics promoted lipid accumulation in HepG2 cells and the additional p53 overexpression rescued the effect of miR-21 (see online supplementary figure S3A, B). We further determined whether p53 deletion could offset the inhibitory effect(s) of miR-21-ASO on lipogenesis. The results showed that miR-21 knockdown led to decreased lipid content and increased p53, and additional treatment with p53 siRNA offset the effects of miR-21-ASO (see online supplementary figure S3C-E). Taken together, our results indicated that miR-21-induced lipid accumulation is, in part, mediated by *HBP1* and p53.

HBP1 mediates the inhibitory effect of miR-21 inhibitor on proliferation, G1/S transition and xenograft tumour from HepG2 cells

To determine whether *HBP1*, at least in part, mediates the effects of miR-21 on proliferation and G1/S transition, we adopted a similar strategy as described above. MTT and soft agar colony assays revealed that miR-21-ASO administration caused suppression of cellular proliferation in HepG2 cells, but additional treatment of *HBP1*-siRNA counteracted the effects of

miR-21-ASO (figure 5D, E). Cell cycle phase distribution of HepG2 cells further showed that miR-21 knockdown led to a significant increase in G1 phase cells and G1/S arrest, but additional treatment of *HBP1*-siRNA antagonised this effect of miR-21-ASO (figure 5F). Consistent with our in vitro findings, miR-21 knockdown inhibited growth of xenograft tumours in nude mice, and *HBP1*-siRNA treatment counteracted the effect of miR-21-ASO (figure 5G).

As we proposed, p53 is an important mediator of the miR-21-Hbp1-p53 axis. To determine whether p53 mediates the inductive effect of miR-21 on proliferation, we treated HepG2 cells with miR-21 mimic or a combination of miR-21 and p53 expression vector. MTT assay, soft agar colony formation assay and cell cycle analysis revealed that miR-21 promoted proliferation and cell cycle progression, and additional overexpression of p53 offset the inductive effects of miR-21 (see online supplementary figure S4A-E). Furthermore, miR-21-ASO treatment led to reduced proliferation, delayed G1/S transition and repressed growth of xenograft tumour, and the additional treatment of p53 siRNA rescued the inhibitory effects of miR-21-ASO (see online supplementary figure S4F-K). In summary, our findings indicated that HBP1 mediates, at least in part, the inhibitory effects of miR-21-ASO on G1/S transition, proliferation and growth of xenograft tumour.



Figure 5 The inhibitory effects of miR-21-anti-sense oligonucleotide (ASO) on lipogenesis, G1/S transition and proliferation are mediated by HBP1. (A and B) miR-21 inhibitor transfection into HepG2 cells reduced lipid content, and additional treatment of HBP1-siRNA restored lipid levels. Lipid droplets were labelled with arrows. Data are presented as mean±SD. Student's t test was used for statistical analysis. (C) miR-21 inhibitor treatment induced expression of HBP1 and p53, and additional knockdown of induced HBP1 with its siRNA inhibited expression of HBP1 and p53. Specifically, HepG2 cells were treated with oleate (0.5 mM) to induce miR-21, and then miR-21-ASO was transfected into HepG2 cells to knock down upregulated miR-21. Levels of miR-21, HBP1 and p53 were determined by qRT-PCR. Data are presented as mean±SD. Student's t test was used for statistical analysis. (D) MTT assay revealed that antagonising miR-21 via miR-21 inhibitor caused reduced cellular proliferation in HepG2 cells, and additional knockdown of HBP1 by its siRNA rescued the effect of miR-21 inhibitor. Data are presented as mean±SD. Student's t test was used for statistical analysis. (E) Soft agar colony formation assay revealed that miR-21 knockdown inhibited the growth of HepG2 cells, and the additional treatment of HBP1-siRNA antagonised the effect of miR-21 inhibitor. Data are presented as mean±SD. Student's t test was used for statistical analysis. (F) miR-21-ASO treatment increased the number of cells in the G1 phase but decreased the number of cells in the S phase, and additional knockdown of upregulated HBP1 by its siRNA antagonised the effect of miR-21 inhibitor. Quantification of the cell cycle phase distribution was analysed by flow cytometry. The proliferation index was reduced in the miR-21 inhibitor treated HepG2 cells and the additional treatment of *HBP*-siRNA offset the effect of miR-21 inhibitor. Data are presented as mean±SD (*p<0.05, **p<0.001, χ^2 test). (G) miR-21-ASO inhibited subcutaneous tumours from HepG2 cells in nude mice, and additional treatment of HBP1-siRNA counteracted the effect of miR-21-ASO. HepG2 cells treated with miR-21-MM-ASO, miR-21-ASO or a combination of miR-21-ASO and HBP1-siRNA were injected subcutaneously into different groups of nude mice. Data are presented as mean±SD. Student's t test was used for statistical analysis.

Antagonising miR-21 in dietary obese mice prevents hepatic lipid accumulation

Next, we assessed the functional contribution of increased *Hbp1* and *p53* expression to hepatic lipid accumulation by reducing miR-21 expression in dietary obese mice. C57Bl/6 mice, which had been on an HFD, were injected with either miR-21-ASO or miR-21-MM-ASO for 4 weeks. We observed that antagonising miR-21 significantly reduced levels of triglycerides in livers of

HFD-treated animals (figure 6A, B), in contrast to plasma triglyceride levels (figure 6C). On the other hand, miR-21-ASO treatment had no effect on body and liver weight (see online supplementary table S6). As expected, we also observed a 91% reduction of hepatic miR-21 expression in mice that received miR-21-ASO compared with miR-21-MM-ASO and a twofold increase of *Hbp1* and *p53* (figure 6D). Four-week treatment of miR-21-ASO showed a strong inhibitory effect on Figure 6 Antagonising miR-21 prevented hepatic lipid accumulation in high fat diet (HFD)-treated mice. (A and B) miR-21 knockdown inhibited lipid accumulation in livers of HFD-fed mice injected with miR-21-anti-sense oligonucleotide (ASO). Representative images are shown. Lipid droplets in livers are labelled with arrows. Cellular triglyceride content was measured by Oil Red staining and triglyceride content was measured with a triglyceride estimation kit. Data are presented as mean±SD (p<0.0002, Mann-Whitney test). (C) miR-21 knockdown had no effect on plasma triglyceride in HFD-treated mice. Data are presented as mean \pm SD (p \geq 0.1, Mann-Whitney test). (D) miR-21-ASO injection into dietary obese mice resulted in downregulated miR-21 and increased *Hbp1* and *p53* expression. Data are presented as mean±SD. Mann-Whitney test was used for statistical analysis. (E) qRT-PCR revealed that HFD-treated mice with decreased levels of miR-21 also retained reduced expression of Scd1, Fasn and Gpat after miR-21-ASO injection. C57Bl/6 mice at 8 weeks of age were kept on HFD for an additional 8 weeks. At 12 weeks of age, mice received a dose of 25 mg/kg miR-21-ASO or miR-21-MM-ASO (0.9% NaCl) weekly for 4 weeks via tail vein injection. Data are presented as mean ±SD. Mann-Whitney test was used for statistical analysis.



hepatosteatosis, but there were no differences in liver and body weight. Therefore, we increased miR-21-ASO treatment time to 8 weeks, which resulted in decreases in both liver weight (see online supplementary table S7) and hepatic lipid content (see online supplementary figure S5B). However, no difference in body weight (see online supplementary table S7), serum free fatty acid and glycerol still was observed (see online supplementary figure S6A, B). In addition, we also observed that miR-21-ASO treatment had no effect on plasma liver enzymes (see online supplementary table S8). These findings indicated that the crosstalk of miR-21 with *Hbp1* and *p53* plays an important role in hepatosteatosis.

We further compared expression levels of *Srebp1c* and lipogenic genes in livers of miR-21-ASO and miR-21-MM-ASO treated mice. As expected, miR-21-ASO treatment led to a significant reduction in *Srebp1c* in the livers of HFD-treated mice (figure 6E). Reduction of *Srebp1c* due to miR-21 knockdown should impair expression of the lipogenic genes. Indeed, in the miR-21-ASO treated group, the mRNA of three enzymes including *Scd1*, *Fasn* and *Gpat* was downregulated at least 1.5 times those of controls (figure 6E).²⁹ Thus, the reduction of *Srebp1c* was associated with a dramatic reduction in the expression of the target enzymes responsible for lipogenesis, which prevented hepatic lipid accumulation. qRT-PCR also revealed that miR-21 knockdown had no effect on β -oxidation-related genes including *Cpt1a*, *Acc2* and *PGC1a* (see online supplementary figure S6C).³² 33

In summary, our data have shown that HFD treatment led to increased miR-21, decreased *Hbp1* and *p53*, which subsequently promoted hepatic lipid accumulation and the potential for HCC, whereas antagonising miR-21 led to the opposite and more therapeutic effect. *Hbp1* is inhibited with increased miR-21 levels and its knockdown impaired transcription of *p53* by *Hbp1*, which led to reduced *p53*. The loss of *p53* then resulted in increased transcription of *Srebp1c*, *CCNB1* and *CCND1*, which promoted both lipogenesis and cell replication (figure 7). Our findings indicate the miR-21 plays an important role in linking NAFLD to HCC by interacting with the *Hbp1-p53-Srebp1c* pathway.

DISCUSSION

Our study addresses a potentially important role for miR-21 in the development of NAFLD and HCC and defines a novel mechanism by which miR-21 contributes to lipogenesis and



Figure 7 Proposed mechanism by which miR-21 links hepatic lipid accumulation and the development of cancer. Hbp1 is a transcriptional activator of *p53*, a suppressor of cell cycle progression and inhibitor of lipogenesis by inhibiting transcription of *Srebp1c*. By directly inhibiting *Hbp1* expression, miR-21 prevents expression of *p53*, which facilitates transcription of genes that are required for lipogenesis and the G1/S transition of cell cycle. As a result, increased miR-21 promotes both hepatic lipid accumulation and potentially carcinogenesis.

carcinogenesis via the *Hbp1-p53-Srebp1c* pathway. The observation that antagonising miR-21 in dietary obese mice potentially improves these metabolic parameters clearly indicates a functional role for increased miR-21 expression in the development of NAFLD. In addition, we also observed that miR-21 knockdown prevented an in situ model of tumorigenesis by targeting *HBP1*. Increased miR-21 expression is not restricted to murine obesity models of NAFLD and HCC,^{19 34} but is also detected in human patients with NASH and HCC.^{6 34 35} Thus, miR-21-ASO may act as a unique potential therapeutic approach for the treatment of both disorders.

Despite its putative role in carcinogenesis,³⁶ the mechanism by which miR-21 regulates NAFLD is unknown. In this study, we observed that HFD administration resulted in increased miR-21 and its knockdown prevented hepatic lipid accumulation. In addition, we have functionally validated Hbp1 as a bona fide target of miR-21, and Hbp1 is a transcriptional activator of p53. It is known that p53 acts as both a tunour suppressor and inhibitor of lipogenesis by inhibiting *Srebp1c*.^{24 25 31} Our findings, combined with those of others, indicate that miR-21 plays critical roles in the pathogenesis of NAFLD and HCC. In addition, miR-21 deletion in p53 knockout mice reduced the incidence of liver cancer;³⁷ and p53 deletion alone promoted hepatosteatosis,²⁵ further supporting the unique role for miR-21 in linking NAFLD to HCC via the Hbp1-p53-Srepb1c pathway. miR-21 significantly inhibits expression of PTEN,³⁴ a negative regulator of NASH in mice,³⁸ suggesting that it might, in part, mediate the inhibitory effect of miR-21 on hepatosteatosis. In this study, we found that Hbp1 is a direct target of miR-21 and confirmed that Hbp1 modulates the inhibitory function of miR-21-ASO on hepatosteatosis and carcinogenesis simultaneously.

We have shown that miR-21 is highly expressed in hepatocytes (figure 1A). Meanwhile, miR-21 is increased but its target Hbp1 is reduced in livers of HFD-treated mice and human individuals with NAFLD and HCC. All these features of miR-21 lead us to focus on its role in linking NAFLD to HCC. Although other miRNAs might contribute to the development of NAFLD and HCC, few other hepatocyte-specific miRNAs meet the above criteria like miR-21. These include increased expression in both NAFLD and HCC of human and mouse, reduced expression of their target genes in mouse models and human patients, and high expression in hepatocytes. We demonstrated that miR-21 knockdown led to upregulated Hbp1 and p53, downregulated Srebp1c and decreased expression of hepatic Scd1, Gpat and Fasn. These findings are consistent with earlier reports that p53 knockout promoted Srebp1c transcription and in turn increased expression of Scd1, Gpat and Fasn, and subsequently hepatic lipogenesis,²⁵²⁸ suggesting that miR-21-dependent Hbp1-p53 pathway inhibition of Srebp1c transcription represents a candidate pathway to cause NAFLD.

There is evidence to suggest that inhibition of p53 attenuates steatosis and liver injury,^{39 40} which is inconsistent with our findings and those of others.^{25 26 41} One possible explanation is that p53 is a negative regulator of hepatic lipid accumulation in the early stages of NAFLD, but p53 is highly expressed at the advanced stages of NAFLD, which may contribute to the high level of apoptosis associated with NASH. In patients with liver steatosis without inflammation, p53 expression was significantly lower than in steatohepatitis,⁴² further suggesting that p53 plays different roles in the various developmental stages of NAFLD. Although the mechanism(s) by which miR-21 controls lipogenesis and carcinogenesis in liver clearly requires further investigation, our study has identified an important role for the interaction of miR-21 with *Hbp1* in obesity-induced NAFLD and HCC. **Contributors** HW and RN: acquisition of data. XC and CJS: analysis and interpretation of data. GS: obtaining funding, study supervision, study concept and design and drafting of the manuscript.

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Competing interests None declared.

Ethics approval All procedures involving mice were approved by the Institutional Animal Care Committee at the University of Minnesota, University of California San Francisco, and the Agency for Science Technology and Research Singapore.

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Supplementary Materials and Methods

Hepatocyte-specific miRNA Profiling

To identify miRNAs that were highly expressed in hepatocytes, we profiled global miRNAs in hepatocytes by comparing miRNA expression of livers between $Dicer1^{fl/fl}$ (n=2) and $Dicer1^{dhep}$ mice (n=2) using Taqman Array MicroRNA Card (Applied Biosystems). These arrays were designed to investigate all miRNAs discovered in human, mouse and rat. Specifically, we isolated total RNA using the miRNeasy kit (Qiagen) from livers of $Dicer1^{fl/fl}$ and $Dicer1^{dhep}$ mice^{1 2}, and the quality of total RNAs were determined using Bioanalyzer 2100. A total of 500 ng of RNA was used for miRNA-specific cDNA synthesis using Megaplex RT Primers and MicroRNA Reverse Transcription Kit (all Applied Biosystems). Taqman MicroRNA Array was run according to the manufacturer's protocol. Data analysis was performed using Viia 7 (Applied Biosystems) and Microsoft Access and the fold change was calculated using $2^{-\Delta\Delta Ct}$ method.³ Internal control was U6 nuclear small RNA and U6 was measured five times in each sample. miR-21 expression was further confirmed between Dicer1 knockout and wild-type mice using Taqman microRNA Assay from Invitrogen.

Fatty Acid Treatment of HepG2 Cells

HepG2 cells were seeded in a 4-well chamber slides using DMEM medium with 10% FBS and allowed to adhere overnight. To determine the inductive effect of miR-21 on lipogenesis, HepG2 cells were then treated with DMEM supplemented with 1% fatty acid free BSA and oleate (0.25 mM). Simultaneously, HepG2 cells cultured in the DMEM containing 0.25 mM oleate were transfected with miR-21 mimic (40 nM) or scrambled control (both Dharmacon) using Lipofectamine 2000. After 24 hours of transfection, Nile-Red Staining was used to

determine the intracellular lipid content in HepG2 cells. To investigate the inhibitory effect of miR-21 inhibitor on lipogenesis, HepG2 cells were then treated with DMEM supplemented with 1% fatty acid free BSA and oleate (0.5 mM). Simultaneously, HepG 2 cells cultured with the DMEM containing 0.5 mM oleate were transfected with miR-21-ASO (40 nM) or miR-21-MM-ASO (control) (Both Exiqon). Lipid accumulation was determined by Nile-Red Staining followed by microfluorimeter detection or imaging.

p53 expression vector (Plasmid #: 12091) and *p53* and *Hbp1* siRNA expression vectors were purchased from Addgene and Origene, respectively; and HBP1 expression vector (pPM-hHBP1-His vector) was purchased from Applied Biological Materials (ABM) Inc. To determine whether the inhibitory effect of miR-21-ASO on lipogenesis is mediated by HBP1, HepG2 cells cultured in the DMEM containing 0.5 mM oleate were transfected with miR-21-ASO (40 nM in 4-well chamber slides), or miR-21-ASO combined with HBP1 siRNA expression vector (200 ng/well in 4-well chamber slides). To determine whether p53 is able to impair the ability of miR-21 mimic to induce lipogenesis, HepG2 cells cultured in the DMEM containing 0.25 mM oleate were transfected with miR-21 mimic (40 nM) or a combination of miR-21 mimic and p53 expression vector (200 ng/ well in 4-well chamber slides). To further determine the role of the interaction between miR-21 and p53, HepG2 cells cultured in the DMEM containing 0.5 mM oleate were transfected with miR-21-ASO (40 nM) or a combination of miR-21-ASO and p53 siRNA (100 ng). Lipofectamine 2000 was used for miR-21 mimic and miR-21-ASO transfection. The cells were cultured for an additional 24 hours, after which lipid accumulation was determined by Nile Red Staining (Sigma-Aldrich).

Lipid Accumulation Assay

The lipid content in HepG2 cells was determined using Nile Red, a vital lipophilic dye (9diethylamino-5H-benzo [alpha] phenoxazine-5-one) from Sigma-Aldrich, which has been shown to selectively stain intracellular lipid droplets. Monolayers were washed with PBS and fixed with 4% paraformaldehyde at room temperature for 10 minutes. After washing, the cells were incubated for 20 minutes with Nile Red solution at a final concentration of 1mg/L in PBS at 37°C. After removal of chamber, the slides were mounted with Prolong® Gold anti-fade reagent with DAPI (Invitrogen) for visualization under fluorescence microscope.

Reporter Vector Construction and Luciferase Assay

To generate the luciferase reporter vectors, *Hbp1* 3' UTR was amplified by PCR from mouse cDNA, and inserted into the pMiR-Reporter vector (Ambion), referred as to pMiR-*Hbp1*. Two bases of the binding site for miR-21 within the 3'UTR of *Hbp1* were mutated using QuikChange II Site-Directed Mutagenesis Kit (Agilent Technologies) per the manufacture's instruction, and referred as to pMiR-*Hbp1-Mu*. Twenty-four hours before transfection, 5×10^4 Hepa1,6 cells were plated per well in a 24-well plate. Then, 200 ng of pMiR-*Hbp1* and miR-21 mimic or miR-21 inhibitor (20 nM or 40 nM) as well as 30 ng of β -*gal* plasmid pSV- β -Galactosidase Control Vector (Promega) were transfected using Lipofectamine 2000 (Invitrogen). Scrambled control (Dharmacon) or miR-21-MM-ASO (Exiqon) was used as the control for miR-21 mimic or miR-21-ASO, respectively.

p53 promoter (1.5 kb) was amplified from mouse genomic DNA and was cloned into pGL3basic vector (Promega), referred as to pGL3-*p53*. 24 hours before transfection, 5×10^4 cells Hepa 1,6 were plated per well in a 24-well plate. Then, 200 ng of pGL3-*p53* with 100 ng or 200 ng *HBP1* expression vector (pPM-hHBP1-His vector) or mice *HBP1* siRNA expression vector (Origene) as well as 30 ng of β-*gal* plasmid pSV-β-Galactosidase Control Vector (Promega) were transfected into Hepa1, 6 cells using Lipofectamine 2000 (Invitrogen). After 24 hours of transfection, luciferase and β -galactosidase assays were done using the Luciferase Assay System and Beta-Glo[®] Assay System (Promega). Luciferase activities were normalized to galactosidase activities; wells were transfected in triplicate; and each well was assayed in triplicate.

Srebp1c promoters (1.5 kb) were amplified from mouse genomic DNA and was cloned into pGL3-basic vector, referred as to pGL3-*Srebp1c*. 24 hours before transfection, 5×10^4 Hepa 1,6 cells were plated per well in a 24-well plate. Then, 200 ng of pGL3-*Srebp1c* with 100 ng or 200 ng *p53* expression vector or *p53* siRNA expression vector (100 ng) and 30 ng of β-*gal* plasmid pSV-β-Galactosidase Control Vector (Promega) were transfected into Hepa1,6 cells using Lipofectamine 2000 (Invitrogen). After 24 hours, luciferase and β-galactosidase assays were performed using the Luciferase Assay System and Beta-Glo[®] Assay System (Promega). Luciferase activities were normalized to galactosidase activities; wells were transfected in triplicate; and each well was assayed in triplicate.

miRNA Transfection and Gene Expression

 5×10^4 of HepG2 cells were seeded in a 24-well plate and allowed to adhere overnight. To determine the effects of miR-21 overexpression and knockdown on gene expression, HepG2 cells cultured in the DMEM with 10% FBS were transfected with miR-21 mimic (40 nM) or inhibitor (40 nM) using Lipofectamine 2000. The equal concentration of scrambled control or miR-21-MM-ASO was used as control for miR-21 mimic or miR-21-ASO, respectively. 24 hours after transfection, cells were washed using cold PBS and the total RNA were isolated for gene expression analysis.

 5×10^4 of HepG2 cells were seeded in a 24-well plate and allowed to adhere overnight. To overexpress *HBP1* or *p53*, 200 ng of pPM-hHBP1-His vector (Applied Biological Materials

(ABM) Inc.,) or *p53* expression vector (Addgene) was transfected into HepG2 cells using Lipofectamine 2000. The control HepG2 cells received empty pcDNATM3.1 vector (Life Technologies). 48 hours after transfection, cells were harvested for RNA isolation and gene expression analysis. To knock down *p53* or *HBP1*, human *p53* siRNA expression vector (100 ng) or *HBP1* siRNA expression vector (100 ng) was introduced into HepG2 cells using Lipofectamine 2000. 48 hours after transfection, the HepG2 cells were collected for RNA isolation and gene expression analysis.

To study whether *HBP1* mediates the inhibitory effect of miR-21-ASO on lipogenesis, 5×10^4 of HepG2 cells were seeded in a 24-well plate and allowed to adhere overnight. Then, the cells cultured in the DMEM containing 0.5 mM oleate were transfected with miR-21-ASO (40 nM), or a combination of miR-21-ASO with *HBP1* siRNA expression vector (200 ng/well). After 48 h, the HepG2 cells were collected for lipid content determination and gene expression analysis. Lipid content in HepG2 cells were determined using Nile-Read staining and gene expression was measured using qRT-PCR.

Histological analysis

Liver samples were embedded in Tissue-Tek OCT embedding compound, and frozen on dry ice. 8 µm-thick sections were cut with a Leica CM3050 S cryostat, air-dried, and fixed in 10% formalin. After washing, sections were stained with an Oil-Red-O (Sigma-Aldrich)/60% isopropanol solution (Fisher Scientific). Briefly, sections were rinsed with 60% isopropanol and stained for 20 min with prepared Oil Red O solution (0.5% in isopropanol followed by dilution to 60% with distilled water and filtered). After rinses in 60% isopropanol and distilled water, slides were counterstained with hematoxylin for 4 min, rinsed with water, and mounted. Hematoxylin and Eosin Staining kit (Scytek laboratories, Inc.) was used in paraformaldehyde-

fixed, paraffin-embedded sections of liver according to manufacturer's protocol. Images were taken with Zeiss Axioplan 2 Upright Microscope.

RNA Isolation and Quantitative Reverse Transcription-PCR (qRT-PCR)

Total RNA was isolated with miRNeasy Mini Kit (Qiagen). To assess gene expression, 1 μ g RNA was used for cDNA synthesis with Superscript III reverse transcription reagent (Invitrogen). PCR amplification was performed at 50°C for 2 minutes and 95°C for 10 minutes, followed by 40 cycles at 95°C for 15 seconds and 60°C for 1 minute in a 7900 real time-PCR system with SYBR green (Applied Biosystems). For each sample, we analyzed β -actin, GAPDH or 18S rRNA expression to normalize target gene expression. Primers for qRT-PCR were designed with Primer Express software (Applied Biosystems). Primers used for quantitative RT-PCR were listed in Supplementary Table 4.

To determine levels of miRNA expression, 10 ng RNA were used for miRNA-specific cDNA synthesis with the TaqMan MicroRNA Reverse Transcription Kit and Taqman MicroRNA Assays (all Applied Biosystems). PCR amplification was performed at 95°C for 10 minutes, followed by 40 cycles at 95°C for 15 seconds and 60°C for 1 minute in a 7900 real time-PCR system (Applied Biosystems). The small RNA Sno202 and RNU6 were used to normalize target miRNA expression. Relative changes in gene and miRNA expression were determined using the $2^{-\Delta\Delta Ct}$ method.³

MTT (3-(4,5-dimethylthiazol-2-yl)-2,5-diphenyltetrazoliumbromide) Assay

Cell proliferation was determined using a MTT cell proliferation kit (Cell Biolabs, Inc.). 5×10^3 of HepG2 cells were seeded in each 96-well plate and allowed to adhere overnight. The cells were then transfected with scrambled control (40 nM), miR-21-ASO (40 nM), or miR-21

mimics (40 nM). After 48 hours culture, cells were used for MTT assay per the manufacturer's instruction (Cell Biolabs, Inc). To determine whether *HBP1* or *p53* mediates the inhibitory effect of miR-21-ASO on cell proliferation, we transfected HepG2 cells with miR-21-ASO (40 nM) or a combination of miR-21-ASO and *HBP1*siRNA expression vector (100 ng) or *p53* siRNA expression vector (100 ng). After 48 hours culture, cells were used for MTT assay per the manufacturer's instruction (Cell Biolabs, Inc). To determine whether *p53* overexpression was able to antagonize the effect of miR-21 on proliferation, we transfected miR-21 mimics (20 nM) or a combination of miR-21 mimics (20 nM) and *p53* expression vector (100 ng) in a 24-well plate. After 48 hours of transfection, cells were used for MTT assay per the manufacturer's instruction (Cell Biolabs, Inc). HepG2 cells were used for MTT assay per the manufacturer's instruction (Cell Biolabs, Inc).

Soft Agar Colony Formation Assay

HepG2 cells (0.5×10^6) in 35-mm plastic dishes were transfected with miR-21 mimics (40 nM) or inhibitors (40 nM). Two days after transfection, transfected cells were suspended with 8 ml of 0.4% top agar (Sigma-Aldrich) and 2×DMEM supplemented with 20% fetal bovine serum before being poured onto 6-cm tissue culture dishes coated with 3.5 ml of 0.7% bottom agar. Fourteen days later, three areas per plate were chosen randomly, the number of visible colonies was counted. To determine whether *HBP1* and *p53* mediate the inhibitory effects of miR-21-ASO on HepG2 cells colony formation, HepG2 cells were transfected with miR-21-ASO (40 nM) or a combination of miR-21-ASO (40 nM) and *p53* siRNA expression vector (100 ng) or *HBP1* siRNA expression vector (1 µg). After transfection for 48 hours, cells were suspended with 8 ml of 0.4% top agar (Sigma-Aldrich) and 2×DMEM supplemented with 20% fetal bovine serum before being poured onto 6-cm tissue culture dishes coated with 3.5 ml of 0.7% bottom agar.

Fourteen days later, three areas per plate were chosen randomly, the number of visible colonies was counted.

Xenograft Tumor Assay

HepG2 cells were placed in a 6-well plate 24 hours prior to transfection. HepG2 cells were transfected miR-21-ASO (40 nM), miR-21-MM-ASO (40 nM) or a combination of miR-21-ASO and *HBP1* siRNA expression vector (1 μ g) or *p53* siRNA expression vector (400 ng). After 24 hours, 5×10⁵ cells in 0.1 ml PBS were injected subcutaneously into the right flank of athymic nude mice (n=9) to establish a model of tumor-bearing mice. Tumor growth was observed every 3 days by measuring its diameter with Vernier calipers. Tumor weight was calculated by gram. Tumor volume (cm³) = d² × D/2, where d is the shortest and D is the longest diameter, respectively. Mice were sacrificed when the tumor size reached 1.5 cm in diameter. All protocols complied with, and all animals received humane care according to, the criteria outlined in the NIH "Guide for the Care and Use of Laboratory Animals."

Cell Cycle Analysis

HepG2 cells were plated in a 6-well plate 24 hours before transfection. After 48 hours of transfection with miR-21 mimic (40 nM) or inhibitor (40 nM), the cells were detached from the plates by trypsin incubation, rinsed with PBS and fixed in 70% (v/v) ethanol. They were then resuspended in PBS and incubated with RNase (100 μ g/ml) and propidium iodide (60 μ g/ml) (Sigma-Aldrich). Cells were analyzed using the FACSCalibur System (BD Biosciences), and the cell cycle phase was analyzed by using CellQuest software. The proliferation index (PI) was calculated as follows: PI = (S+G2/M)/G1. S, G2/M and G1 refer to the percentage of cells in S phase, G2/M phase and G1 phase, respectively. PI=(S+G2/M)/G1. To determine whether *p53* or *HBP1* mediates the inhibitory effect of miR-21-ASO on cell cycle progression, HepG2 cells

were transfected with miR-21-ASO (40 nM) or a combination of miR-21-ASO (40 nM) and *HBP1* siRNA expression vector (1 μ g) or *p53* siRNA expression vector (100 ng). After 48 hours of transfection, the HepG2 cells were treated as above for cell cycle analysis.

Human Cell Cycle RT² ProfilerTM PCR Array

We determined the effect of *HBP1* on expression of genes controlling cell cycle using Human Cell Cycle RT² *Profiler*TM PCR Array (Qiagen). The Human Cell Cycle RT² *Profiler*TM PCR Array profiles the expression of 84 genes key to cell cycle regulation. Briefly, HepG2 cells were plated in a 6 well plate 24 hours prior to transfection. HepG2 cells were transfected with 1 µg of pPM-hHBP1-His vector (Applied Biological Materials (ABM) Inc.,) using Lipofectamine 2000. The control HepG2 cells were treated with empty pcDNATM3.1 vector (Life Technologies). 48 hours after transfection, cells were harvested for RNA extract. A total of 2 µg RNA was used to perform the reverse transcription using SuperScript® III Reverse Transcriptase (Invitrogen). Cell Cycle RT² *Profiler*TM PCR Array was run according to the manufacturer's protocol. Data analysis was performed using $2^{-\Delta\Delta Ct}$ method.³ Internal control is β -actin. *p53* expression was further confirmed between *HBP1* expressed vector treated cells and pcDNA3.1 vector treated cells using qRT-PCR.

Plasma Lipid Analysis

Blood was collected into tubes, containing 4 mM of EDTA, from cardiac puncture of C57Bl/6 mice after 4 or 8 weeks of HFD treatment. Plasma was separated by centrifugation (3000 x RPM for 20 min at 4 °C and triglyceride (mg/dl, Roche Diagnostics) was quantified enzymatically. Serum chemistry was carried out by the Pathology Laboratory of the University of Minnesota.

Hepatic Lipid Analysis

Mouse liver (100 mg) was placed in 1 ml chloroform/methanol (2:1) mixture and incubated on mice for 10 minutes before homogenization. Lipids were extracted from liver homogenates through room temperature orbital shaking (2 hours) followed by centrifugation (5000 RPM for 5 minutes). Supernatants were collected and washed with 0.4 ml chloroform/methanol (2:1) mixture by centrifugation at 5000 RPM for 20 minutes (room temperature). New supernatants were washed with 0.2 volume of 0.9% NaCl. After centrifuging for 5 minutes at 5000 RPM, supernatants were removed and lower-phase was dried at 42°C. Dried lipids were re-suspended in 2% Triton X-100. Liver triglycerides were quantified via a colorimetric assay using a triglyceride assay kit from Roche Diagnostics according to the manufacturer's protocols.

Western Blot Analysis

Western blot was performed following standard procedures. HBP1 primary antibodies were purchased from Abcam; and binding was visualized using SuperSignal west femto maximum sensitivity substrate (Cat # 34095, Thermo Fisher Scientific Inc.).

References

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- 2. Mattis AN, Song G, Hitchner K, Kim RY, Lee AY, Sharma AD, et al. A screen in mice uncovers repression of lipoprotein lipase by microRNA-29a as a mechanism for lipid distribution away from the liver. *Hepatology* 2015; 61(1):141-52.
- 3. Schmittgen TD, Livak KJ. Analyzing real-time PCR data by the comparative C(T) method. *Nature Protoc* 2008;3(6):1101-8.

Supplementary Figures:

Figure 1. HFD treatment increases both hepatic and plasma triglycerides. (A) Oil-Red staining showing that HFD treatment led to increased hepatic lipid accumulation in mice. Lipid droplets in livers were labeled with arrows. (B,C) HFD treatment led to high levels of hepatic and plasma triglycerides. Lipids were extracted from the livers of mice treated with standard diet (n=6) or HFD (n=6) and triglycerides measured using a colorimetric assay. Representative histological images are shown. Data are expressed as mean \pm SD. Mann-Whitney was used for statistical analysis. (D) HFD treatment led to reduced expression of *Hbp1* in livers of dietary obese mice. Data are expressed as mean \pm SD. Mann-Whitney was used for statistical analysis.



Figure 2. (A) Overexpression of p53 caused dose-dependent inhibition of the activity of a luciferase reporter gene linked to the *Srebp1c* promoter. (B) p53 knockdown via its siRNA released the inhibitory effect of p53 on *Srebp1c* transcription, which was reflected by increased luciferase activity. Data are expressed as mean \pm SD. In this multiple-groups experiment, we only performed comparison between two groups and Student T test was used for statistical analysis.



Supplementary Figure 2

Figure 3. miR-21 modulates lipid accumulation in HepG2 cells by interacting with *p53*. (A, B) Transfection of miR-21 mimic into HepG2 cells incubated with oleate led to increased intracellular lipid content, and additional overexpression of *p53* offset the effect of miR-21. Lipid droplets in human hepatocytes were labeled with arrows. Data are expressed as mean \pm SD. Student T test was used for statistical analysis. (C,D) miR-21 inhibitor transfection into HepG2 cells cultured with the medium containing 0.5 mM oleate led to a decrease in intracellular lipid content, and additional treatment of *p53* siRNA antagonized the effect of miR-21 inhibitor on lipid accumulation. Lipid droplets in human hepatocytes were labeled with arrows. Data are expressed as mean \pm SD. Student T test was used for statistical analysis. (E) Mechanically, miR-21 inhibitor treatment induced expression of *p53* and additional transfection of *p53* siRNA into HepG2 cells knocked down induced *p53*. Briefly, HepG2 cells were cultured with DMEM containing 0.5 mM oleate. 24 hours after miR-21 inhibitor transfection, *p53* siRNA was further introduced into HepG2 cells in attempt to knock down *p53* induced by miR-21 inhibitor. Data are expressed as mean \pm SD. Student T test was used for statistical analysis.



Supplementary Figure 3

Figure 4. The inhibitory effects of miR-21-ASO on cell cycle progression and proliferation are mediated by p53. (A) miR-21 mimic transfection into HepG2 cells inhibited expression of p53, and treatment of p53 expression vector significantly increased expression of p53. Data are presented as mean ± SD. Student T test was used for statistical analysis. (B) MTT assay revealed that miR-21 overexpression induced proliferation, and overexpression of p53 counteracted the inductive effect of miR-21 on proliferation. Data are presented as mean \pm SD. Student T test was used for statistical analysis. (C) Soft agar colony formation assay revealed that miR-21 overexpression promoted the colony formation of HepG2 cells, and further overexpression of p53antagonized the effect of miR-21. Data are presented as mean \pm SD. Student T test was used for statistical analysis. (D,E) miR-21 mimic treatment led to a decrease in the number of cells in the G1 phase but an increase in the number of cells in S phase, and additional overexpression of p53antagonized the effect of miR-21. Quantification of the cell cycle phase distribution was analyzed by flow cytometry. The proliferation index was increased in the miR-21 mimic-treated HepG2 cells and overexpression of p53 offset the effect of miR-21. Data are presented as mean \pm SD (*p < 0.05, **p < 0.001, Chi-Square Test). (F) miR-21 inhibitor transfection into HepG2 cells induced expression of p53, and additional knockdown of induced p53 with its siRNA inhibited expression of p53. Specifically, HepG2 cells were treated with oleate (0.5 mM) to induce miR-21, and then miR-21-ASO was transfected into HepG2 cells to knock down upregulated miR-21. Levels of p53 were determined by qRT-PCR. Data are presented as mean \pm SD. Student T test was used for statistical analysis. (G) MTT assay revealed that antagonizing miR-21 via miR-21 inhibitor led to reduced cellular proliferation in HepG2 cells, and additional knockdown of p53 by its siRNA rescued the effect of miR-21 inhibitor. Data are presented as mean \pm SD. Student T test was used for statistical analysis. (H) Soft agar colony formation assay revealed that miR-21 knockdown inhibited the growth of HepG2 cells, and the additional treatment of p53-siRNA antagonized the effect of miR-21 inhibitor. Data are presented as mean \pm SD. Student T test was used for statistical analysis. (I,J) miR-21 inhibitor treatment prevented cell cycle progression, and additional knockdown of p53 rescued the effect of miR-21 inhibitor. miR-21 knockdown increased the number of cells in the G1 phase but decreased the number of cells in S phase, and additional knockdown of up-regulated p53 by its siRNA antagonized the effect of miR-21 inhibitor. Quantification of the cell cycle phase distribution was analyzed by flow cytometry. The proliferation index was reduced in the miR-21 inhibitor treated HepG2 cells and the additional treatment of p53-siRNA offset the effect of miR-21 inhibitor. Data are presented as mean \pm SD (**p < 0.001, Chi-Square Test). (K) miR-21-ASO inhibited growth of subcutaneous tumors from HepG2 cells in nude mice, and additional treatment of p53-siRNA counteracted the effect of miR-21-ASO. HepG2 cells treated with scramble control, miR-21-ASO or a combination of miR-21-ASO and p53-siRNA were injected subcutaneously into nude mice. Data represent mean \pm SD. Mann-Whitney was used for statistical analysis of tumor weight.

Supplementary Figure 4



Figure 5. Knocking down miR-21 prevented hepatic lipid accumulation in HFD-treated mice. (A,B) miR-21 knockdown inhibited lipid accumulation in livers of HFD-fed mice injected with miR-21-ASO. Representative images (Oil-Red) are shown. Lipid droplets in livers were labeled with arrows. Cellular triglyceride content was measured by Oil Red staining and triglyceride (TG) content (per mg protein) was measured with a triglyceride estimation kit. Briefly, 8-week old mice were kept on HFD until 20 weeks of age. At 12 weeks of age, mice were divided into two groups: one group (n=8) received miR-21-MM-ASO and the other group received miR-21-ASO for another eight weeks. (C) miR-21-ASO injection into dietary obese mice resulted in down-regulated miR-21 and increased *Hbp1* and *p53* expression. Data represent mean \pm SD. Mann-Whitney was used for statistical analysis. **p* < 0.05.

miR-21-MM-ASO miR-21-ASO Α **Oil-Red** в С 200 2.5 miR-21 p = 0.0002 Hbp1 160 2 Liver triglycerides mg/g Fold change p53 120 1.5 80 1 40 0.5 0 0 Control miR-21 Control miR-21-ASO -ASO

Supplementary Figure 5

Figure 6. miR-21 knockdown had no effect on serum free fatty acid and glycerol as well as the expression of the genes encoding enzymes for fatty acid oxidation. (A,B) miR-21-ASO treatment had no effect on levels of serum free fatty acid and glycerol in dietary obese mice. (C) qRT-PCR revealed that miR-21 knockdown did not change the expression of β -oxidation-related genes including *Cpt1a*, *Acc2* and *PGC1a*.



Supplementary Table 1. 1219 down-regulated probes in livers of human patients with NAFLD/NASH.

Probe ID	p-value	Log (Fold Change)	Gene Symbol	Gene Description
7916225	0.00002042	-0.79297091	RPS13	ribosomal protein S13
8095744	0.00003301	-0.92809751	AREG	amphiregulin
8016739	0.00003413	-0.76219365	TOB1	transducer of ERBB2, 1
8121850	0.00007819	-0.96944624	HEY2	hairy/enhancer-of-split related with YRPW motif 2
8166219	0.00013805	-0.52888646	SYAP1	synapse associated protein 1
7907970	0.00017988	-0.63720968		
7924058	0.00019095	-0.62327473	IRF6	interferon regulatory factor 6
8152642	0.00019118	-0.58996711	FAM83A-AS1	FAM83A antisense RNA 1
7892578	0.00020598	-0.86107507		
8167356	0.00021234	-0.5746145	GLOD5	glyoxalase domain containing 5
8122732	0.00021362	-0.69869308		
8115831	0.00024035	-1.39904664	DUSP1	dual specificity phosphatase 1
8174361	0.00024065	-1.10270028	TSC22D3	TSC22 domain family, member 3
7910901	0.00025567	-0.57259484	LOC100130331	POTE ankyrin domain family, member F pseudogene
7980096	0.00026905	-1.06351259		
8104136	0.00031457	-0.57617112	HMX1	H6 family homeobox 1
7899018	0.00034108	-0.58214279	TMEM57	transmembrane protein 57
8154381	0.00040117	-0.89537625	LURAP1L	leucine rich adaptor protein 1-like
8044574	0.00042422	-1.31166294	IL1RN	interleukin 1 receptor antagonist
7893573	0.00042537	-0.8413454		
7894916	0.00043289	-0.74484068		
7966631	0.00044308	-0.52391017	LHX5	LIM homeobox 5
8180222	0.00045126	-0.73639191	CFHR4	complement factor H-related 4
7908499	0.00045842	-1.02950046	CFHR5	complement factor H-related 5
8132013	0.00050245	-0.41798903	CHN2	chimerin 2
8144982	0.00055067	-0.47603714	NPM2	nucleophosmin/nucleoplasmin 2
8081036	0.00056767	-0.86673648	CADM2	cell adhesion molecule 2
8123800	0.00058719	-0.53901395		
8098782	0.00065132	-0.52548035	CPLX1	complexin 1
7946504	0.00068011	-0.45301645	TMEM41B	transmembrane protein 41B
8121794	0.00076598	-0.74999714	SMPDL3A	sphingomyelin phosphodiesterase, acid-like 3A
8149809	0.00077964	-0.42161283		
8069574	0.00083873	-0.6975465	C21orf91	chromosome 21 open reading frame 91
8040898	0.00084825	-0.60386321	TRIM54	tripartite motif containing 54
7989611	0.00088558	-0.64984027	FAM96A	family with sequence similarity 96, member A
8024485	0.00088896	-1.31050838	GADD45B	growth arrest and DNA-damage-inducible, beta
8180275	0.00089652	-0.73017018	PCDHA1	protocadherin alpha 1
7893001	0.00091425	-0.61391617		
8167625	0.00097516	-0.43744251	CCNB3	cyclin B3
8122317	0.00098027	-0.40045385	HEBP2	heme binding protein 2
8094938	0.00099488	-0.78036905	NIPAL1	NIPA-like domain containing 1
7961371	0.00103814	-0.61238977	DUSP16	dual specificity phosphatase 16
8111136	0.00106935	-1.05109382	FAM134B	family with sequence similarity 134, member B
8034698	0.00110117	-1.02876002	MIR23A	microRNA 23a
8140386	0.0011019	-0.4358513	STYXL1	serine/threonine/tyrosine interacting-like 1
8137526	0.00110482	-0.82985909	INSIG1	insulin induced gene 1
8174675	0.00120137	-0.53338897	SLC25A5-	SLC25A5 antisense RNA 1///solute carrier family 25 (mitochondrial carrier;
8024020	0.00122152	0 42402007	AS1///SLC25A5	adenine nucleotide transiocator), member 5
7804060	0.00133133	-0.42402097	A201	
1094200	0.00137202	-0.00404844		transmin T type 1 (alcolated, alcow)
0039340	0.00151127	-0.43677351		uoponini i type i (skeletal, siow)
0000330	0.00153113	-1.04839709		Cysteme-senne-non nuclear protein 1
0152215	0.00158019	-0.5133994		Nupper-like factor 10///Krupper-like factor 10
7928218	0.0015906	-0.79662233		caunenn-related 23
7947199	0.00161768	-0.46886289		LUDD N activitative protein coupled receptor 4
1906863	0.00162373	-0.44543897		DDF-IN-acteyigiucosamine pyrophosphorylase 1
8052934	0.00163613	-0.47586977	WUCEE	
8040362	0.00168743	-0.69510683		

7967789	0.00174682	-0.64920848	PXMP2	peroxisomal membrane protein 2, 22kDa
7895162	0.00178057	-0.46039958		
8083569	0.00178308	-0.79711107	TIPARP	TCDD-inducible poly(ADP-ribose) polymerase
7892628	0.00181803	-0.72775738		
7892797	0.00181805	-0.43085863		
7895866	0.00183265	-0.96302911		
8033956	0.00183473	-0.41030632	S1PR2	sphingosine-1-phosphate receptor 2
7969177	0.00185368	-0.43959976	ST13P4	suppression of tumorigenicity 13 (colon carcinoma) (Hsp70 interacting
				protein) pseudogene 4
8025828	0.00185487	-1.15153203	LDLR	low density lipoprotein receptor
7959827	0.00187837	-0.35015972	TMEM132C	transmembrane protein 132C
7985898	0.00194622	-0.37701062	WDR93	WD repeat domain 93
7962537	0.00195513	-0.98323651	SLC38A2	solute carrier family 38, member 2
8158554	0.00197004	-0.37942865	PRRX2	paired related homeobox 2
8096077	0.00202598	-1.01498157		
8096661	0.00206275	-0.4885318		
8032273	0.00210311	-0.48392927		
8126946	0.00212526	-0.42875452		
8028652	0.00216135	-1.21798455	ZFP36	ZFP36 ring finger protein
8173106	0.00219743	-0.4609312	ITIH6	inter-alpha-trypsin inhibitor heavy chain family, member 6
8067955	0.00220035	-0.4353128	CXADR	coxsackie virus and adenovirus receptor
7927814	0.0022399	-0.47780987	SIRT1	sirtuin 1
8036902	0.00227133	-0.56879519	SERTAD1	SERTA domain containing 1
8034286	0.00228385	-0.43253	ECSIT	ECSIT signalling integrator
8169352	0.00229803	-0.58505298	NXT2	nuclear transport factor 2-like export factor 2
8138741	0.00229978	-0.45670343	HOXA6	homeobox A6
7922162	0.00231424	-0.8041639	SLC19A2	solute carrier family 19 (thiamine transporter), member 2
7997257	0.00233541	-0.55018592	ZFP1	ZFP1 zinc finger protein
7943218	0.00239565	-0.4273094	PANX1	pannexin 1
8101648	0.00240798	-0.6334724	HSD17B11	hydroxysteroid (17-beta) dehydrogenase 11
7938368	0.00241197	-0.58274175		
7928630	0.00241664	-0.37087115	EIF5AL1///EIF5A	eukaryotic translation initiation factor 5A-like 1///eukaryotic translation
7005400	0.00040000	0.00007005		initiation factor 5A
7925128	0.00242339	-0.62687995	PPP00	
7925813	0.0024389	-0.56368497		Drolle rich 26
7987099	0.00248494	-0.44641745		
7974870	0.00248683	-0.52323329	SNAPCT	
8095341	0.00258274	-0.51313788	04 = #(50///04 = #(50	
7905324	0.00259819	-0.50365441	C10ff56///C10ff56	chromosome 1 open reading frame 56///chromosome 1 open reading frame 56
7958000	0.00260001	-0.41980653		choline phosphotransferase 1
8040473	0.00260512	-0.71029488		ras homolog family member B
8027345	0.00260822	-0.44951545	ZNF492	zinc finger protein 492
7894489	0.00262948	-0.43569551		
8126474	0.00269744	-0.3809945	PPP2R5D///MEA1	protein phosphatase 2, regulatory subunit B', delta///male-enhanced antigen 1
/940971	0.00269815	-0.476535	KUNK4///TEX40	potassium channel, subtamily K, member 4///testis expressed 40
8156309	0.00280094	-1.91857805	GADD45G	growth arrest and DNA-damage-inducible, gamma
8052123	0.00282848	-0.49124312		
8114185	0.0028402	-0.36110419	CDKN2AIPNL	CDKN2A interacting protein N-terminal like
8025515	0.00287786	-0.44084668	RDH8	retinol dehydrogenase 8 (all-trans)
8167862	0.00289701	-0.48092759	PAGE2B	P antigen family, member 2B
8055287	0.00289919	-0.43796413	MZ12A	mitotic spindle organizing protein 2A
8043743	0.00292461	-0.67967357		
8174026	0.00293023	-0.55828121	YWHAQP8	YWHAQ pseudogene 8
8152355	0.00296149	-0.77105697	SYBU	syntabulin (syntaxin-interacting)
8042107	0.00297762	-0.50763377	EIF3F	eukaryotic translation initiation factor 3, subunit F
8103524	0.00304423	-0.32197155	TMEM192	transmembrane protein 192
8044927	0.00306715	-0.73908427	INHBB	inhibin, beta B
7893298	0.00307572	-1.04787723		
7902282	0.00307667	-0.38643169	HHLA3	HERV-H LTR-associating 3
7955896	0.00308094	-0.46793182	COPZ1	coatomer protein complex, subunit zeta 1
8018482	0.00308408	-0.46854697	WBP2	WW domain binding protein 2
7987381	0.00308548	-0.37073497	CSNK1A1P1	casein kinase 1, alpha 1 pseudogene 1
8097655	0.00312283	-0.56275697		

7933084	0.00319926	-0.90824229	NAMPT	nicotinamide phosphoribosyltransferase
7899932	0.00321168	-0.39531876	GJB3	gap junction protein, beta 3, 31kDa
7984008	0.00322888	-0.60391894		
7938348	0.00323796	-0.71731105	WEE1	WEE1 homolog (S. pombe)
8109663	0.00327684	-0.37097365	GABRA1	gamma-aminobutyric acid (GABA) A receptor, alpha 1
8037535	0.00334481	-0.5266372		
7931348	0.00338259	-0.41154445	FOXI2	forkhead box I2
8057599	0.00339997	-0.49814204	TFPI	tissue factor pathway inhibitor (lipoprotein-associated coagulation inhibitor)
7972062	0.00341863	-0.53169338	FBXL3	F-box and leucine-rich repeat protein 3
7922523	0.00341999	-0.72352437		
8084035	0.00342392	-0.3806076	ZNF639	zinc finger protein 639
8002556	0.00343743	-0.41156863	ТАТ	tyrosine aminotransferase
8077944	0.00356907	-0.44030569	CAND2	cullin-associated and neddylation-dissociated 2 (putative)
8048319	0.00358257	-1.34572837	VII 1	villin 1
8030999	0.00360323	-0.73399042	ZNF331	zinc finger protein 331
7893809	0.00363236	-0.48877502		
7939492	0.00363515	-0 59049103	C11orf96	chromosome 11 open reading frame 96
8086028	0.00372278	-0.36696877	GLB1	calactosidase beta 1
8120562	0.0037/056	-1 10524489	CTGE	connective tissue arouth factor
7007655	0.00374950	0.50/2288/	0101	
8020507	0.00380401	-0.39433004		poliovirus receptor related 2 (boroesvirus entry mediator B)
0029507	0.00301324	-0.40000473		form 1 homolog o (C. clogopo)
8124001	0.00390139	-0.44379653		
8134091	0.00397943	-0.39300923		ciduulli 12
8030299	0.00400619	-0.36263339	CCDC 155	
8165907	0.00401462	-0.42457406	00//740	a sete de serve a se s'al ser e sede con 't VIII a contra con t'al s o (l'e se s)
8127629	0.00412227	-0.43334274		cytochrome c oxidase subunit vila polypeptide 2 (liver)
7937330	0.00412576	-0.72633257	IFIIM2	interferon induced transmembrane protein 2
8161884	0.00413807	-0.3905697	PRUNE2	prune homolog 2 (Drosophila)
8040103	0.00418069	-0.50310353	ID2	inhibitor of DNA binding 2, dominant negative helix-loop-helix protein
7933723	0.00418145	-0.48001218	IPMK	inositol polyphosphate multikinase
7977340	0.00421513	-0.61037656	BTBD6	BTB (POZ) domain containing 6
8070665	0.00422801	-2.17844415	SIK1	salt-inducible kinase 1
8150592	0.00424845	-0.92584694	CEBPD	CCAAT/enhancer binding protein (C/EBP), delta
7931553	0.00425034	-0.44916502	UTF1	undifferentiated embryonic cell transcription factor 1
7906819	0.00426254	-0.42172089	ATF6	activating transcription factor 6
8028563	0.0043084	-0.36296876	SARS2///MRPS12	seryl-tRNA synthetase 2, mitochondrial///mitochondrial ribosomal protein S12
7968272	0.00434427	-0.40411607		
8124848	0.00435247	-1.24562714	IER3	immediate early response 3
7955589	0.00438037	-1.83708367	NR4A1	nuclear receptor subfamily 4, group A, member 1
8075332	0.00440358	-0.34419627	TBC1D10A	TBC1 domain family, member 10A
7926531	0.00440758	-0.78966531	ARL5B	ADP-ribosylation factor-like 5B
8172028	0.00443478	-0.56427871		
7963061	0.00445042	-0.39782438	C1QL4	complement component 1, q subcomponent-like 4
8076734	0.00446229	-0.32904051	WNT7B	wingless-type MMTV integration site family, member 7B
7997245	0.00447818	-0.57646424		
8022912	0.0044798	-0.42757287	MIR187	microRNA 187
7928823	0.00450568	-0.46092822	OPN4	opsin 4
7896542	0.00452104	-0.44880106		
8029489	0.00457471	-0.58404465	BCAM	basal cell adhesion molecule (Lutheran blood group)
8041561	0.0045827	-0.69651727	MORN2	MORN repeat containing 2
8056102	0.00458612	-0.47832061	LY75-	LY75-CD302 readthrough///CD302 molecule///CD302 molecule
			CD302///CD302///C D302	
8172654	0.00460281	-0.4908284	USP27X-AS1	USP27X antisense RNA 1 (head to head)
8089011	0.00460815	-0.58678728	PROS1	protein S (alpha)
8032051	0.00462119	-0.48997936	ODF3L2	outer dense fiber of sperm tails 3-like 2
8026341	0.00462468	-0.39211721	MIR639///TECR	microRNA 639///trans-2,3-enoyl-CoA reductase
7979906	0.00462887	-0.33595199	COX16	COX16 cytochrome c oxidase assembly homolog (S. cerevisiae)
8067585	0.00463215	-0.48589407	BHLHE23	basic helix-loop-helix family, member e23
8169080	0.00464251	-0.46899293	H2BFM	H2B histone family, member M
8055952	0.00467622	-2.38162242	NR4A2	nuclear receptor subfamily 4. group A. member 2
8145440	0.00468513	-0.4207451	PPP2R2A	protein phosphatase 2, regulatory subunit B, alpha
				, , , , , , , , , , , , , , , , , , ,

7997281	0.00474822	-0.31522401	LOC100288358///TE RF2IP	uncharacterized LOC100288358///telomeric repeat binding factor 2, interacting p
7921473	0.00475568	-0.40804966	CCDC19	coiled-coil domain containing 19
8001658	0.00475659	-0.49442371	C16orf80	chromosome 16 open reading frame 80
7984011	0.00479953	-0.4501947	FOXB1	forkhead box B1
7966321	0.00481561	-0.48311361	GPN3	GPN-loop GTPase 3
8081548	0.00491062	-0.34885596	PVRL3	poliovirus receptor-related 3
7961702	0.00493373	-0.51273359	KCNJ8	potassium inwardly-rectifying channel, subfamily J, member 8
7902290	0.00497914	-0.85723533	CTH	cystathionase (cystathionine gamma-lyase)
7893126	0.00499271	-0.50921745		
7895999	0.00508128	-0.80143545		
8038477	0.00513034	-0.39106372	AKT1S1	AKT1 substrate 1 (proline-rich)
7992447	0.00515645	-0.46435071	SYNGR3	synaptogyrin 3
7988033	0.00515705	-0.34336504	EPB42	erythrocyte membrane protein band 4.2
8042211	0.00518106	-0.40948862	B3GNT2	UDP-GlcNAc:betaGal beta-1,3-N-acetylglucosaminyltransferase 2
8175775	0.00525884	-0.36505992	MAGEA1	melanoma antigen family A, 1 (directs expression of antigen MZ2-E)
8113220	0.00528051	-0.4084689	ELL2	elongation factor, RNA polymerase II, 2
8041048	0.00528818	-0.76435982	FOSL2	FOS-like antigen 2
8027390	0.0053856	-0.57256643	POP4	processing of precursor 4, ribonuclease P/MRP subunit (S. cerevisiae)
7910638	0.00539462	-0.32394432		
7971905	0.00541491	-0.99462736	PCDH20	protocadherin 20
8173613	0.00544019	-0.47524059	RLIM	ring finger protein, LIM domain interacting
7904921	0.00548926	-0.41853402		
8075198	0.00552375	-0.38715775		
8149475	0.0055617	-0.40361036	CNOT7	CCR4-NOT transcription complex, subunit 7
7896417	0.00563782	-0.94617864		
7912361	0.0056567	-0.36774972	MASP2	mannan-binding lectin serine peptidase 2
8022404	0.0056903	-0.49151104	FAM210A	family with sequence similarity 210, member A
8173174	0.00570365	-0.44169143	USP51	ubiquitin specific peptidase 51
8019273	0.00572308	-0.4437143	ALYREF	Aly/REF export factor
8062782	0.00574316	-0.35107174	TOX2	TOX high mobility group box family member 2
8152092	0.00574696	-0.49306521		
7908496	0.00576171	-0.30036417	CFHR2	complement factor H-related 2
7935679	0.00584077	-0.59294195	CPN1	carboxypeptidase N, polypeptide 1
7898448	0.00586683	-0.32563272	PADI4	peptidyl arginine deiminase, type IV
7903474	0.00588765	-0.66252786		
8088167	0.00590185	-0.73221561	SELK	selenoprotein K
8053386	0.00593987	-0.42835305		
7896709	0.00595633	-0.54491053		
7977621	0.00596052	-0.31595099	NDRG2	NDRG family member 2
7970241	0.00599773	-0.47770509	F10	coagulation factor X
7930413	0.00605122	-0.59449389	DUSP5	dual specificity phosphatase 5
7895018	0.0060528	-0.76877776		
7904084	0.00607682	-0.42246545	AKR7A2P1	aldo-keto reductase family 7, member A2 pseudogene 1
8134452	0.00610284	-0.3608268	BHLHA15	basic helix-loop-helix family, member a15
8012891	0.00614412	-0.44763079		
7979886	0.00615054	-0.47851425		
7972745	0.00617206	-0.99279147	IRS2	insulin receptor substrate 2
7945688	0.00619404	-0.32391	INS- IGF2///INS///IGF2	INS-IGF2 readthrough///insulin///insulin-like growth factor 2 (somatomedin A)
8080960	0.00621401	-0.40331914		
8131475	0.00623208	-0.42100182	C1GALT1	core 1 synthase, glycoprotein-N-acetylgalactosamine 3-beta-galactosyltransfera
8035398	0.00623283	-0.60831376	RAB3A	RAB3A, member RAS oncogene family
7993185	0.00627444	-0.45327951	NUBP1///NUBP1	nucleotide binding protein 1///nucleotide binding protein 1
7978666	0.00630918	-0.40811759	MBIP	MAP3K12 binding inhibitory protein 1
8123763	0.00635391	-0.32196566		
8106976	0.00640806	-0.38056911	GPR150	G protein-coupled receptor 150
7905533	0.00646517	-0.42610209	IVL	involucrin
8082667	0.00652538	-0.42223065	NUDT16	nudix (nucleoside diphosphate linked moiety X)-type motif 16
7911138	0.00656876	-0.32565054	CNST	consortin, connexin sorting protein
8172022	0.00662795	-0.39992152	TMEM47	transmembrane protein 47
8061507	0.00665163	-0.39298354		
7905949	0.0067186	-0.41477061		

8172345	0.00671888	-0.3772574	ELK1	ELK1, member of ETS oncogene family
7898655	0.00672387	-0.60517746	CDA	cytidine deaminase
8132694	0.00672768	-2.42071729	IGFBP1	insulin-like growth factor binding protein 1
8135480	0.00675373	-0.58554821	DNAJB9	DnaJ (Hsp40) homolog, subfamily B, member 9
8141533	0.00678251	-0.40249095		
8111757	0.00680358	-0.47485865	C9	complement component 9
7991809	0.00681807	-0.43076631	PDIA2///ARHGDIG	protein disulfide isomerase family A, member 2///Rho GDP dissociation inhibitor
8052087	0.00687575	-0.46199213		
8043682	0.00687815	-0.59561521	LOC653924	glycerol-3-phosphate acyltransferase 2, mitochondrial pseudogene
8157264	0.00689371	-0.59903063	SLC31A2	solute carrier family 31 (copper transporters), member 2
7909142	0.00693518	-0.53564924	NUCKS1	nuclear casein kinase and cyclin-dependent kinase substrate 1
8019181	0.0069406	-0.4619242	LOC100130370	uncharacterized LOC100130370
8012344	0.00697218	-0.45144554	HES7	hairy and enhancer of split 7 (Drosophila)
7921014	0.00703173	-0.37471617	MEF2D///MEF2D	myocyte enhancer factor 2D///myocyte enhancer factor 2D
7894011	0.00705212	-0.64433672		
7920697	0.00708493	-0.38311175	GBA	glucosidase, beta, acid
8037737	0.00711773	-0.32678056	NOVA2	neuro-oncological ventral antigen 2
8036031	0.00717089	-0.47881442		
8023984	0.0071809	-0.3533444	HCN2	hyperpolarization activated cyclic nucleotide-gated potassium channel 2
8082605	0.00721481	-0.34669716		
7931556	0.00723762	-0.42116855	VENTX	VENT homeobox
7917741	0.00724415	-0.35159811	TMED5	transmembrane emp24 protein transport domain containing 5
8112613	0.00725973	-0.55011642		
8006562	0.00728147	-0.44878483	RASL10B	RAS-like, family 10, member B
8034514	0.00734294	-0.30934348	C19orf43	chromosome 19 open reading frame 43
7909610	0.00742968	-0.80696302	ATF3	activating transcription factor 3
8059026	0.00743657	-0.33410252	MIR375	microRNA 375
8014865	0.00748496	-0.30968903	NEUROD2	neuronal differentiation 2
7902687	0.00749592	-1.58929102	CYR61	cysteine-rich, angiogenic inducer, 61
8129666	0.00752051	-0.65162755	SLC2A12	solute carrier family 2 (facilitated glucose transporter), member 12
8079140	0.00753724	-0.41811603	SNRK	SNF related kinase
7929653	0.00753755	-0.41250631	ANKRD2	ankyrin repeat domain 2 (stretch responsive muscle)
7952408	0.00758054	-0.51254783	SIAE	sialic acid acetylesterase
7893679	0.00762347	-0.66557408		
8012000	0.00765775	-0.5342722	RNASEK	ribonuclease, RNase K
7951660	0.00766067	-0.31945717		
8101699	0.00767203	-0.76333399	PPM1K	protein phosphatase, Mg2+/Mn2+ dependent, 1K
8073960	0.0076739	-0.34498536	PIM3	pim-3 oncogene
8029693	0.00776588	-2.48951161	FOSB///FOSB	FBJ murine osteosarcoma viral oncogene homolog B///FBJ murine osteosarcom
8125010	0.00778632	-0 70085020		nomolog B
0123313	0.00770032	-0.70003929	5	ancharacterized E00200047777 N300 binding protein 3
8124433	0.00780294	-0.39079902	HIST1H4G	histone cluster 1, H4g
7920047	0.00782009	-0.30044311	MRPL9	mitochondrial ribosomal protein L9
8005747	0.00782995	-0.40879915		
8070257	0.00784738	-0.42553747	PIGP	phosphatidylinositol glycan anchor biosynthesis, class P
8042335	0.00785852	-0.39836679	VDAC2	voltage-dependent anion channel 2
7896657	0.00790549	-0.36538957		
8063386	0.00791892	-0.45700192	CEBPB	CCAAT/enhancer binding protein (C/EBP), beta
7938748	0.00792251	-0.45622772		
7912207	0.00793853	-0.41925277		
7931926	0.00797927	-0.30257615		
8112570	0.00798736	-0.31578424	MRPS27	mitochondrial ribosomal protein S27
8099918	0.00808893	-0.39758671		
8099633	0.0080994	-0.93030989	PPARGC1A	peroxisome proliferator-activated receptor gamma, coactivator 1 alpha
7892679	0.00817767	-0.7803035		
8123463	0.00825871	-0.40715449	C6orf120///PHF10	chromosome 6 open reading frame 120///PHD finger protein 10
8168892	0.00828467	-0.30980017	TCEAL2	transcription elongation factor A (SII)-like 2
7895253	0.00832644	-0.7513525		
8164428	0.00833208	-0.31929364	COQ4///TRUB2	coenzyme Q4 homolog (S. cerevisiae)///TruB pseudouridine (psi) synthase hom
7896321	0.00834717	-0.63173595		
8098604	0.00836097	-1.0878421	ANKRD37///UFSP2	ankyrin repeat domain 37///UFM1-specific peptidase 2
8161610	0.00853768	-0.62339209	TMEM252	transmembrane protein 252

8146559	0.00854129	-0.4153998		
8013804	0.00855972	-0.46431045	DHRS13	dehydrogenase/reductase (SDR family) member 13
8079117	0.00856495	-0.70857495	CCBP2	chemokine binding protein 2
8038877	0.00857639	-0.50946137	SIGLEC5	sialic acid binding Ig-like lectin 5
8134415	0.0085817	-0.34670333	ACN9	ACN9 homolog (S. cerevisiae)
7934278	0.00861048	-1.41272271	P4HA1	prolyl 4-hydroxylase, alpha polypeptide I
8105151	0.00862206	-0.36808791		
7908488	0.00863954	-0.43377221	CFHR1	complement factor H-related 1
7895972	0.00864048	-0.57482767		
8149289	0.00865386	-0 43339143	SOX7	SRY (sex determining region Y)-box 7
8109732	0.0086566	-0.40228854	MAT2B	methionine adenosyltransferase II beta
8116658	0.00866762	-0.37468245	FAM50B	family with sequence similarity 50 member B
8096489	0.00000702	-0.3052260		PDZ and LIM domain 5
7070448	0.000000040	-0.31183144	G IB6	r b2 and Ein domain 3
810440	0.000704	0.37805016		gap Junction protein, beta 0, 30kDa
7026220	0.00877865	-0.37893910		ribosomal protoin 1122 psoudogono 6
7930320	0.00077803	-0.43326203		late endeemel/wassemel adapter MARK and MTOR activator 5
7916437	0.00002001	-0.34163604	LAIMTORS	ale endosomal/lysosomal adaptor, MAPK and MITOR activator 5
7936833	0.0088747	-0.64274924		
7895263	0.0088885	-0.38760549	N#4050414	
8133114	0.00889094	-0.32720993		vitamin K epoxide reductase complex, subunit 1-like 1
8104022	0.0089315	-0.59880096	PDLIM3	PDZ and LIM domain 3
/944006	0.00893599	-0.48927117	KBM/	KNA binding motif protein /
8166382	0.00897159	-0.31889815	MBTPS2	membrane-bound transcription factor peptidase, site 2
7966389	0.00897588	-0.33924654	FAM109A	family with sequence similarity 109, member A
7995324	0.00897682	-0.58131206		
7945536	0.00897846	-0.4207714	CEND1	cell cycle exit and neuronal differentiation 1
7910790	0.00898445	-0.5331333		
7937465	0.00899656	-0.33036562	TALDO1	transaldolase 1
8053139	0.00903744	-0.32251651	C2orf81	chromosome 2 open reading frame 81
8115144	0.00903823	-0.40536169	ARSI	arylsulfatase family, member I
7985777	0.00904812	-0.35551268	ISG20	interferon stimulated exonuclease gene 20kDa
7919568	0.00905407	-0.44556348		
7965152	0.009072	-0.40563482		
8027402	0.00912961	-0.36855138	CCNE1	cyclin E1
8096533	0.00914464	-0.41241511		
8147566	0.00914732	-0.37727663	KCNS2	potassium voltage-gated channel, delayed-rectifier, subfamily S, member 2
7950644	0.00915194	-0.31693204	NDUFC2	NADH dehydrogenase (ubiquinone) 1, subcomplex unknown, 2, 14.5kDa
				methylonetetrohydrofolete dehydrogenege (NADD) denendent) 1. methenyltetro
7975045	0.00915666	-0.40716591	MTHFD1	I melliylenelelianyulululale uenyuluqenase (NADF+ uepenueni) 1, melhenyilelia
7975045	0.00915666	-0.40716591	MTHFD1	cyclohydrolase, formyltetrahydrofolate synthetase
7975045 7995332	0.00915666 0.00920364	-0.40716591 -0.55891548	MTHFD1	cyclohydrolase, formyltetrahydrofolate synthetase
7975045 7995332 7940005	0.00915666 0.00920364 0.00924711	-0.40716591 -0.55891548 -0.47656773	MTHFD1 P2RX3	cyclohydrolase, formyltetrahydrofolate synthetase purinergic receptor P2X, ligand-gated ion channel, 3
7975045 7995332 7940005 8070689	0.00915666 0.00920364 0.00924711 0.00929776	-0.40716591 -0.55891548 -0.47656773 -0.372955	MTHFD1 P2RX3 HSF2BP	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein
7975045 7995332 7940005 8070689 8060741	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199	MTHFD1 P2RX3 HSF2BP	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein
7975045 7995332 7940005 8070689 8060741 8117696	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342	MTHFD1 P2RX3 HSF2BP COX11	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast)
7975045 7995332 7940005 8070689 8060741 8117696 7947828	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.0094803	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.0094803 0.00949417	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.0094503 0.00949417 0.00951594	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.0094503 0.00949417 0.00951594 0.00953039	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRE2P1	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.0094503 0.00949417 0.00951594 0.00953039 0.00954886	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.4444967 -0.48114118	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.0094503 0.00951594 0.00953039 0.00954886 0.00955327	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.4444967 -0.48114118 -0.42479366	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.0094803 0.00949417 0.00951594 0.00953039 0.00954886 0.0095503	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.4444967 -0.48114118 -0.42479366 -0.58428935	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1 flavin containing monooxygenase 5
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7000397	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.0094803 0.00949417 0.00951594 0.00953039 0.0095503 0.0095503 0.0095503 0.00955774	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731022	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1 flavin containing monooxygenase 5
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895194	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.00945475 0.00945475 0.00951594 0.0095503 0.0095503 0.0095503 0.00957671 0.00957671	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59026259	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2	interfiger letter any diologie deny diogenase (NADP + dependent) 1, methely tetra cyclohydrolase, formyltetrahydrofolate synthetase purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1 flavin containing monooxygenase 5 epithelial membrane protein 2
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895181	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.00945475 0.00945475 0.00951594 0.0095503 0.0095503 0.0095503 0.00957671 0.00961533 0.00963234	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59036358	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2	<pre>interinterinterinterinterinterinterinter</pre>
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895181 8146837 769555	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.00945475 0.00945475 0.00951594 0.0095503 0.0095503 0.0095503 0.00957671 0.00961533 0.00963234	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59036358 -0.57234307 -0.42092	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2	purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 chromosome 9 open reading frame 41 flavin containing monooxygenase 5 epithelial membrane protein 2
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895181 8146837 7935855 7095322	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.00945475 0.00945475 0.00951594 0.00951594 0.0095503 0.00955503 0.00957671 0.00961533 0.00963234 0.00970267 0.0097027	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59036358 -0.57234307 -0.32919828 -0.2248022	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2 LBX1	Purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 chromosome 9 open reading frame 41 flavin containing monooxygenase 5 epithelial membrane protein 2
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895181 8146837 7995326 7995326	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.00945475 0.00945475 0.00951594 0.0095503 0.00955503 0.00955503 0.00955503 0.00957671 0.00961533 0.00963234 0.00970578 0.00970578	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59036358 -0.57234307 -0.32919828 -0.39318939 -0.59144755	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2 LBX1	<pre>interfigure for the form of the form</pre>
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895181 8146837 7995326 7995326 7917240	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.0094803 0.00949417 0.00951594 0.00953039 0.0095503 0.00955503 0.00955503 0.00955503 0.00957671 0.00961533 0.00963234 0.00970278 0.00972789	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59036358 -0.57234307 -0.32919828 -0.39318939 -0.57141755 -0.4020112	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2 LBX1 CTBS	Interfyterfeterfarlyddiolate derfyddiogenase (NADP + dependent) 1, methefyterfa cyclohydrolase, formyltetrahydrofolate synthetase purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1 flavin containing monooxygenase 5 epithelial membrane protein 2 ladybird homeobox 1 chitobiase, di-N-acetyl-
7975045 7995332 7940005 8070689 8060741 8117696 7947828 8106473 8161829 8094874 8016320 8044682 8132209 7922754 7919314 7948369 7999387 7895181 8146837 7995326 7995326 7917240 7999171	0.00915666 0.00920364 0.00924711 0.00929776 0.00934063 0.00934483 0.00938372 0.00938475 0.00945475 0.00945475 0.00945475 0.00945039 0.0095503 0.0095503 0.00955503 0.00955503 0.00957671 0.00955503 0.00957671 0.00961533 0.00963234 0.00970289 0.00972789 0.00977018	-0.40716591 -0.55891548 -0.47656773 -0.372955 -0.35615199 -0.49963342 -0.40675709 -0.43605402 -0.43605402 -0.43462103 -0.39952773 -0.44895129 -0.44605661 -0.4444967 -0.48114118 -0.42479366 -0.58428935 -0.35731933 -0.59036358 -0.57234307 -0.32919828 -0.39318939 -0.57141755 -0.46860413 -0.565757	MTHFD1 P2RX3 HSF2BP COX11 MYBPC3 C9orf41 RPRML ZNRF2P1 FMO5 EMP2 LBX1 CTBS	<pre>interlyterleterarlyterlotate derlytotgerlase (NADF+ dependent) 1, metherlyterla cyclohydrolase, formyltetrahydrofolate synthetase purinergic receptor P2X, ligand-gated ion channel, 3 heat shock transcription factor 2 binding protein cytochrome c oxidase assembly homolog 11 (yeast) myosin binding protein C, cardiac chromosome 9 open reading frame 41 reprimo-like zinc and ring finger 2 pseudogene 1 flavin containing monooxygenase 5 epithelial membrane protein 2 ladybird homeobox 1 chitobiase, di-N-acetyl-</pre>

7894504	0.0098639	-0.81397023		
8150830	0.00986768	-0.47849016	LYPLA1	lysophospholipase I
8056943	0.00991212	-0.35405226	KIAA1715	KIAA1715
8098414	0.00993869	-0.56395089	SPCS3	signal peptidase complex subunit 3 homolog (S. cerevisiae)
8110604	0.00997563	-0.44555367		
8161238	0.01003471	-0.50454921	RAB1C	RAB1C, member RAS oncogene family pseudogene
8076655	0.01014197	-0.32891458	PNPLA5	patatin-like phospholipase domain containing 5
7980990	0.01014993	-0.55239339	C14orf142///UBR7	chromosome 14 open reading frame 142///ubiquitin protein ligase E3 componen (putative)
8147970	0.01015589	-0.30581758	EBAG9	estrogen receptor binding site associated, antigen, 9
8088908	0.01020996	-0.41134802		
7938756	0.01025138	-0.32092892	ST13P5	suppression of tumorigenicity 13 (colon carcinoma) (Hsp70 interacting protein) p
8106475	0.0103189	-0.3594368	ACTB	actin, beta
7977951	0.01036025	-0.34461725		
8148304	0.01039175	-1.18304529	TRIB1	tribbles homolog 1 (Drosophila)
7939902	0.01040063	-0.38828584	LOC646813	DEAH (Asp-Glu-Ala-His) box helicase 9 pseudogene
8097801	0.01042103	-0.72860338	FAM160A1	family with sequence similarity 160, member A1
8146118	0.01042813	-0.45186442		
8105104	0.0104285	-0.44920844	C5orf51	chromosome 5 open reading frame 51
8096457	0.01048316	-0.31630198		
8170882	0.01048448	-0.30836951	ATP6AP1	ATPase, H+ transporting, lysosomal accessory protein 1
7950471	0.01048504	-0.31612176	OR2AT4	olfactory receptor, family 2, subfamily AT, member 4
8120937	0.01053038	-0.31732211	RIPPLY2	ripply2 homolog (zebrafish)
8035435	0.01053185	-0.41787212	KIAA1683	KIAA1683
8117071	0.01053456	-0.38635547	FAM8A1	family with sequence similarity 8, member A1
8005473	0.01053554	-0.61759142	PAIP1	poly(A) binding protein interacting protein 1
7979725	0.01064551	-0.6167953	PLEKHH1///PIGH	pleckstrin homology domain containing, family H (with MyTH4 domain) member 1///phosphatidylinositol glycan anchor biosynthesis, class H
7895045	0.0106535	-0.3972872		
7895382	0.01067106	-0.43690162		
7959946	0.01067747	-0.36204361	MMP17	matrix metallopeptidase 17 (membrane-inserted)
8055862	0.01067783	-0.52379533	ARL5A	ADP-ribosylation factor-like 5A
8045075	0.01067908	-0.3331529	GPR17	G protein-coupled receptor 17
8149330	0.01076585	-0.32968311	CTSB	cathepsin B
8016433	0.01077936	-0.35821508	HOXB1	homeobox B1
8167165	0.01081851	-0.36529185	ARAF	v-raf murine sarcoma 3611 viral oncogene homolog
8133219	0.01082924	-0.34597685		
8037301	0.01085019	-0.36746609	LYPD3	LY6/PLAUR domain containing 3
7975779	0.01094785	-2.88929943	FOS	FBJ murine osteosarcoma viral oncogene homolog
8169709	0.01097268	-0.59147705	GLRX5	glutaredoxin 5
8103411	0.01107409	-0.51160094		
8169233	0.01107828	-0.89697503		
8159583	0.01108292	-0.30872508	GRIN1	glutamate receptor, ionotropic, N-methyl D-aspartate 1
7917599	0.01115141	-0.38245791	BARHL2	BarH-like homeobox 2
8002051	0.01122705	-0.4209698	AGRP	agouti related protein homolog (mouse)
8015273	0.01124124	-0.47729307	KRI31	keratin 31
7993126	0.0112/186	-0.35256905	ABAT	4-aminoputyrate aminotransterase
8090872	0.01130951	-0.3586311	KY MDU0	kypnoscollosis peptidase
8133690	0.01133111	-0.33750963		malate denydrogenase 2, NAD (mitochondrial)
8007471	0.01133115	-0.34387362		neignbor of BRCA1 gene 1
7972808	0.01140737	-0.45470767		carbonydrate kinase domain containing
8025375	0.01141417	-0.33462212	CCL25	chemokine (C-C motif) ligand 25
8020455	0.01142591	-0.32615/6/		GATA binding protein b
7970498	0.01143682	-0.32320218	LAISZ	arge turnor suppressor kinase 2
7909700	0.01148112	-0.38455019	GAIC	
1893539	0.01148932	-0.89483126	ClartEl	abromosomo 2 open reading frame 50
7004044	0.01149917	-0.010/2419	0301136	chromosome s open reading frame so
7076024	0.011506/2	-0.02109/82	MIR404	microPNA 404
1910834	0.01109/4/	-0.33291097		IIIUUUNNA 494
0100403	0.01162163	-0.37204271		solute carrier raminy 4, sourium bicarbonate corransporter, member 9
0109240	0.01160000	-0.47039001		prospriorituosyl pyropriospriate synthetase 1
0039273	0.01109896	-0.30604803		CDC42 enector protein (Kno G Pase pinding) 5
1900381	0.01172681	-0.78031434	EFCAB4B	EF-nand calcium binding domain 4B

7906469	0.01172837	-0.43181692	DUSP23	dual specificity phosphatase 23
8007208	0.01176499	-0.37747576	HSPB9	heat shock protein, alpha-crystallin-related, B9
8039504	0.01185048	-0.32685578	SHISA7	shisa homolog 7 (Xenopus laevis)
8172803	0.01185087	-0.35944288		
8127425	0.01187957	-0.33122812	LMBRD1	LMBR1 domain containing 1
8147687	0.01188765	-0.96720356		•
8103706	0.0119034	-0.43042592	AADAT	aminoadipate aminotransferase
8151436	0.01191743	-0.40114317	PEX2	peroxisomal biogenesis factor 2
8037197	0.01199463	-0.35368453	CXCL17	chemokine (C-X-C motif) ligand 17
8148317	0.01213641	-1.53771242	MYC	v-myc myelocytomatosis viral oncogene homolog (avian)
7995007	0.01214603	-0.44895844	HSD3B7	hydroxy-delta-5-steroid dehydrogenase, 3 beta- and steroid delta-isomerase 7
8091485	0.01215067	-0.46352355	SIAH2	siah E3 ubiquitin protein ligase 2
8055992	0.01221026	-0.44396258	ACVR1C	activin A receptor, type IC
8013259	0.01225764	-0.37267665		
8045321	0.01226119	-0.34637321	POTEF///POTEE	POTE ankyrin domain family, member F///POTE ankyrin domain family, membe
7906978	0.01226135	-0.39136656	MGST3///MGST3	microsomal glutathione S-transferase 3///microsomal glutathione S-transferase 3
7989473	0.01226217	-0.46402123	C2CD4B	C2 calcium-dependent domain containing 4B
8175811	0.01230799	-0.38985992	FAM58A///FAM58A	family with sequence similarity 58, member A///family with sequence similarity 5
8073192	0.01233734	-0.33962969	UQCRFS1	ubiquinol-cytochrome c reductase, Rieske iron-sulfur polypeptide 1
8164883	0.01234507	-0.35384973	SURF4	surfeit 4
8129039	0.01241569	-0.43250449		
7901445	0.01248125	-0.34184383		
7937106	0.01249689	-0.40268783	NKX6-2	NK6 homeobox 2
8162624	0.01253841	-0.80813545	AAED1	AhpC/TSA antioxidant enzyme domain containing 1
7970096	0.01256054	-0.30243621	ING1	inhibitor of growth family, member 1
7931500	0.01258157	-0.35851563	GPR123	G protein-coupled receptor 123
8040949	0.01260771	-0.49050292	KRTCAP3	keratinocyte associated protein 3
7949995	0.01263317	-0.37894332	MRPL21	mitochondrial ribosomal protein L21
8070900	0.01265558	-0.3728143		
8088820	0.01270739	-0.39483186	RYBP	RING1 and YY1 binding protein
7929947	0.01280099	-0.3864066	TLX1	T-cell leukemia homeobox 1
8024623	0.01280121	-0.36112096	NFIC	nuclear factor I/C (CCAAT-binding transcription factor)
7949894	0.01280713	-0.38023445		
7893236	0.01281341	-0.46729035		
7972973	0.01282642	-0.31533021	SETD8	SET domain containing (lysine methyltransferase) 8
8108050	0.01286117	-0.38233943	TCF7	transcription factor 7 (T-cell specific, HMG-box)
7909225	0.0128639	-0.39847863	DYRK3	dual-specificity tyrosine-(Y)-phosphorylation regulated kinase 3
8052676	0.01291357	-0.35226701		
8113413	0.01297933	-0.44238759	NUDT12	nudix (nucleoside diphosphate linked moiety X)-type motif 12
8067201	0.01299057	-0.4433533		
8059965	0.01299385	-0.32936562	KLHL30-AS1	KLHL30 antisense RNA 1
8012349	0.01301791	-0.61080484	PER1	period circadian clock 1
8001108	0.01307271	-0.45434015		
7950128	0.01307978	-0.40380926	LRTOMT///ANAPC1	leucine rich transmembrane and 0-methyltransferase domain containing///anaph
70000	0.04040=	0.5000.1=1=	5	complex subunit 15
7922646	0.01310747	-0.52034567	TOR1AIP2	torsin A interacting protein 2
8107468	0.0131189	-0.43346861		
7896094	0.01311953	-0.46778485		
8162276	0.01318523	-1.27354794	NFIL3	nuclear factor, interleukin 3 regulated
/931810	0.01321711	-1.08633018	KLF6	Kruppei-like factor 6
/893850	0.01322589	-0.61197901	TUDDOD	
8123651	0.01326686	-0.34933522	TUBB2B	tubulin, beta 2B class lib
7908525	0.01329628	-0.3526/747	C10ff53	chromosome 1 open reading frame 53
7938100	0.01338751	-0.40413009		spningomyelin phosphodiesterase 1, acid lysosomal
8149685	0.0134127	-0.31159078	LGI3	leucine-rich repeat LGI family, member 3
7924910	0.0134202	-0.33775024	ACTA1	actin, alpha 1, skeletal muscle
/980001	0.01352574	-0.42469421	MDDOC	
8159249	0.01354097	-0.38676713	MRPS2	mitochondrial ribosomal protein S2
7933209	0.01354222	-0.302417	C100ff25///RASSF4/	cnromosome 10 open reading trame 25///Ras association (RalGDS/AF-6) doma
7995322	0.01356053	-0.50750183		
8153409	0.01361469	-0.43716573	MAFA	v-mat musculoaponeurotic fibrosarcoma oncogene bomolog A (avian)
7898721	0.01365553	-0.39126966		
1000121	0.01000000	0.00120000		

7978093	0.01365727	-0.36393444	JPH4	junctophilin 4
7989708	0.01371619	-0.3938201	MTFMT	mitochondrial methionyl-tRNA formyltransferase
8063796	0.01375761	-0.30485353	CDH4	cadherin 4, type 1, R-cadherin (retinal)
7940662	0.01379246	-0.30687787	ROM1	retinal outer segment membrane protein 1
7996331	0.01379648	-0.34469461	CA7	carbonic anhydrase VII
8096281	0.01380044	-0.3148774	IBSP	integrin-binding sialoprotein
7992145	0.01384314	-0.3655483	SSTR5	somatostatin receptor 5
7978838	0.01387614	-0.39360271	DNAAF2	dynein, axonemal, assembly factor 2
7920971	0.01389666	-0.44245375	C1orf85	chromosome 1 open reading frame 85
7934178	0.0139363	-0.3468346	PCBD1	pterin-4 alpha-carbinolamine dehydratase/dimerization cofactor of hepatocyte n
	0.04004400	0.04050054	115054	alpha
8141643	0.01394108	-0.34059854	UFSP1	UFM1-specific peptidase 1 (non-functional)
7893685	0.01395807	-0.36791291	70004	to other a shell a d
7992191	0.01396418	-0.36515952	TPSD1	tryptase delta 1
7893344	0.01398267	-0.41858942	05700	
8145047	0.01398749	-0.35209255	SFIPC	
7944401	0.01399267	-0.3230911	HMBS	nydroxymetnyibilane synthase
8095139	0.01419659	-0.49530176	SRD5A3	steroid 5 alpha-reductase 3
8158147	0.01429382	-1.03344562	SLC25A25	solute carrier family 25 (mitochondrial carrier; phosphate carrier), member 25
7893196	0.01431305	-0.48386921		- this we denote Plan AA
8023920	0.01441801	-0.49887911	TXNL4A	
1892916	0.01442825	-0.39090428		free fatty and recentor 1
8027854	0.01458923	-0.38329854	FFAR1	
7997861	0.01459896	-0.35865965		
8105508	0.0146273	-0.36157331		NOTCH-regulated ankyrin repeat protein
7999718	0.01465169	-0.3490034		
8121727	0.01467365	-0.33059573	BRD7P3	bromodomain containing 7 pseudogene 3
7950267	0.01406366	-0.3676406		Whith inducible signaling potential and protein 2
0002004	0.01470961	-0.32015008	WI5P2	whit i inducible signaling pathway protein 2
0110024	0.01475166	-0.43460547	DDK1	PRICK1 SCARAWAVE acting nucleating complex subunit
0003207	0.0147041	-0.43712020		TAEOR DNA polymoroog II. TATA boy binding protoin (TPD) opposited factor
0170203	0.01400200	-0.31743060	TAF9D	TAP9B RNA polymerase II, TATA box binding protein (TBP)-associated factor, s
0073012	0.01401000	-0.31790703		
0029070	0.01400450	-0.32363693	WASI	Mickett Aldrich oundrome like
8001178	0.01490459	-0.30949337	C16orf87	chromosomo 16 opon roading framo 87
8110920	0.01512884	-0.43219034		
8019046	0.01512004	-0.3651434	FIF4A3	eukanyotic translation initiation factor 443
8124134	0.01517148	-0.30223801		thiopurine S-methyltransferase
780/881	0.0152645	-0.3581052		
7905329	0.0152858	-0.6738789	MUT11	myeloid/lymphoid or mixed-lineage leykemia (trithoray homolog, Drosophila): tra
7896637	0.01537938	-0 39231734		
7903281	0.01539254	-0.41256119	SI C35A3	solute carrier family 35 (UDP-N-acetylolucosamine (UDP-GlcNAc) transporter)
8003425	0.01545441	-0.336718	CBFA2T3	core-binding factor, runt domain, alpha subunit 2: translocated to 3
8158666	0.01547562	-0.49568416	HMCN2	hemicentin 2
7931764	0.01547816	-0.34974081	ADARB2	adenosine deaminase, RNA-specific, B2 (non-functional)
8089112	0.01556802	-0.69116131	FILIP1L	filamin A interacting protein 1-like
8149612	0.01558939	-0.35665093	LZTS1	leucine zipper, putative tumor suppressor 1
7939424	0.01564802	-0.36768455	API5	apoptosis inhibitor 5
7942774	0.01567066	-0.40242264	AQP11	aquaporin 11
8035234	0.01576141	-0.39597498		
8165453	0.01605512	-0.39955056	LRRC26	leucine rich repeat containing 26
7896533	0.01609657	-0.67037838		
7981078	0.01613337	-0.49110393	SERPINA11	serpin peptidase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member
7909332	0.01614287	-0.57211101	CD55	CD55 molecule, decay accelerating factor for complement (Cromer blood group
8172471	0.01618281	-0.35593395	PIM2	pim-2 oncogene
8037267	0.01618553	-0.474422	PSG2	pregnancy specific beta-1-glycoprotein 2
8042701	0.01621342	-0.36762858	EMX1	empty spiracles homeobox 1
8116859	0.01621874	-0.33113292	TMEM14C	transmembrane protein 14C
8024373	0.01626396	-0.32987556	IZUMO4	IZUMO family member 4
8110708	0.01627093	-0.47987818	TPPP	tubulin polymerization promoting protein
8171837	0.01630449	-0.54013801	KLHL15	kelch-like family member 15

7895971	0.01630642	-0.54769836		
8030578	0.01637273	-0.31031437	FLJ26850	FLJ26850 protein
8040249	0.01640558	-0.31062974	ATP6V1C2///PDIA6	ATPase, H+ transporting, lysosomal 42kDa, V1 subunit C2///protein disulfide isc member 6
8055688	0.0164527	-0.83364087	RND3///RND3	Rho family GTPase 3///Rho family GTPase 3
8101992	0.01652567	-0.37992405	SLC39A8	solute carrier family 39 (zinc transporter), member 8
8139421	0.01654418	-0.36508112	H2AFV	H2A histone family, member V
8067167	0.01655575	-0.60701203	AURKA	aurora kinase A
8155508	0.01655871	-0.75226649		
8055672	0.01658109	-0.35416777	MMADHC	methylmalonic aciduria (cobalamin deficiency) cbID type, with homocystinuria
8125843	0.01659299	-0.31380661	SPDEF	SAM pointed domain containing ets transcription factor
7955624	0.01666015	-0.41859108	KRT86///KRT83///K	keratin 86///keratin 83///keratin 81
8035868	0.01668323	-0.304017	RT81	
8160465	0.01000323	-0.304017		
7079644	0.016754931	-0.37509556		nuclear factor of kanna light polynoptide gone ophonoor in P calls inhibitor, alph
7978644	0.01675560	-0.43907969		nuclear factor of kappa light polypeptide gene enhancer in B-cells inhibitor, alph
7956395	0.01675569	-0.37916514		
7895752	0.01680823	-0.55872325	00007	O secto in some had as sector 07
8080958	0.0170633	-0.54267042	GPR27	G protein-coupled receptor 27
7896519	0.01708722	-0.58062185	0111 704 /	
8037949	0.01/1//69	-0.41832376	SULT2A1	sulfotransferase family, cytosolic, 2A, dehydroepiandrosterone (DHEA)-preferrin
8097687	0.01718892	-0.38931962	POU4F2	POU class 4 homeobox 2
8158902	0.01724479	-0.40478951	C9orf171	chromosome 9 open reading frame 171
7981084	0.01724911	-0.35802564	SERPINA9	serpin peptidase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member
7898537	0.01728462	-0.31245777	PAX7	paired box 7
8156240	0.01728631	-0.30924988	CTSL3P	cathepsin L family member 3, pseudogene
8024228	0.0173331	-0.31886935	MIDN	midnolin
7957551	0.0174208	-1.02415093	SOCS2	suppressor of cytokine signaling 2
8110427	0.01744307	-0.32370395		
8110166	0.01744636	-0.41437139	HIGD2A	HIG1 hypoxia inducible domain family, member 2A
8063970	0.01746431	-0.30210449	COL9A3	collagen, type IX, alpha 3
8058866	0.01748739	-0.31716286	TNP1	transition protein 1 (during histone to protamine replacement)
7895902	0.01750909	-0.57093101		
8030706	0.01751678	-0.37022035	EMC10	ER membrane protein complex subunit 10
7981013	0.01757047	-0.31638017	PRIMA1	proline rich membrane anchor 1
8025028	0.01759601	-0.3261772	CLPP	ClpP caseinolytic peptidase, ATP-dependent, proteolytic subunit homolog (E. co
7902930	0.01760699	-0.39332551	HSP90B3P	heat shock protein 90kDa beta (Grp94), member 3, pseudogene
8119102	0.01761731	-0.44535647		
7925161	0.0176747	-0.42738364	IRF2BP2	interferon regulatory factor 2 binding protein 2
7912968	0.0177088	-0.43397222	TAS1R2	taste receptor, type 1, member 2
8027920	0.0177103	-0.32199904	FTV2	ets variant 2
7969975	0.01777618	-0 43521291		
7893026	0.01777863	-0.33833864		
8112422	0.01790304	-0.33107496		
7966746	0.01796364	-0.42326506	HRK	harakiri, BCI 2 interacting protein (contains only BH3 domain)
7978586	0.01802709	-0.5384399	CFL2	cofilin 2 (muscle)
7997158	0.0181073	-0.34428531		MARVEL domain containing 3
8073544	0.01813965	-0.31870155	MIR33A	microRNA 33a
8065602	0.0181/679	-0.35227612		testis specific protein. V-linked 26. pseudogene
8038380	0.01014070	-0.30/222012		rotasome (proceme macronain) 265 subunit non ATDaco 9
8157605	0.01010/91	-0.50430387		protoasonie (prosonie, macropani) 200 suburnt, 1101-ATEase, 0
7900776	0.01010413	-0.0023413		
1032110	0.01019091	-0.4304710	MCC	mutated in colorectal cancers
0113001	0.01019237	-0.40306933		
0130337	0.01821486	-0.42728155		
7900021	0.01822571	-0.89783926	202	senne denydratase
1893339	0.01824485	-0.52709092		
8115664	0.01826251	-0.3860378		giutaredoxin (thioitransferase)
8084275	0.01827394	-0.30541854		5-nydroxytryptamine (serotonin) receptor 3C, ionotropic
8022674	0.01831479	-0.38186196	CDH2	caonerin 2, type 1, N-cadherin (neuronal)
/896378	0.01834395	-0.73322051	75410-	
8161747	0.01838533	-0.79970969	ZFAND5	zinc tinger, AN1-type domain 5
7894827	0.01840224	-0.38072453		
7893962	0.01843417	-0.70487892		

8119620	0.01844262	-1.15973477	GNMT	glycine N-methyltransferase
8027385	0.0184487	-0.449968	VSTM2B	V-set and transmembrane domain containing 2B
8030128	0.01847241	-0.81531956	PPP1R15A	protein phosphatase 1, regulatory subunit 15A
7927146	0.01850802	-0.36234914	CSGALNACT2	chondroitin sulfate N-acetylgalactosaminyltransferase 2
7895096	0.01859161	-0.46518114		
8083826	0.01859301	-0.40926291	SEC62	SEC62 homolog (S. cerevisiae)
7895127	0.01859667	-0.62104955		
7995342	0.01859732	-0.57088018		
8069633	0.01859803	-0.32686666	ATP5J	ATP synthase. H+ transporting, mitochondrial Fo complex, subunit F6
8098240	0.01865334	-0.53738251		
8103023	0.01866578	-0.51929891		
7971345	0.01870527	-0.40222941		
8021484	0.01870848	-0.32247486	CDH20	cadherin 20, type 2
7893624	0.01875201	-0.54683872	02.1.20	
7893458	0.01876127	-0.50923216		
8056728	0.01878338	-0.32988846		
8083136	0.01890196	-0 56952413	ATP1B3	ATPase Na+/K+ transporting beta 3 polypeptide
8037166	0.01894193	-0 32733753	FRE	Ets2 repressor factor
7025342	0.01806050	0.55169033		EBQ1 like beta (S. corovisiao)
7925542	0.01090039	-0.55106955	ERUILD	EROT-like bela (S. Celevisiae)
8005233	0.01902063	-0.03109102		iun Dinrete encorrene
8035445	0.01903955	-0.06262462		Jun D proto-oncogene
8025968	0.01922158	-0.40112981	ZNF69	zinc finger protein 69
8118667	0.01926325	-0.49673901	MYL12B	myosin, light chain 12B, regulatory
8156196	0.01930364	-0.31685026	C90ff170	chromosome 9 open reading frame 170
8104369	0.01938669	-0.55158501	SRD5A1	steroid-5-alpha-reductase, alpha polypeptide 1 (3-oxo-5 alpha-steroid delta 4-de alpha 1)
8050763	0.019394	-0.33115268	PTRHD1	peptidyl-tRNA hydrolase domain containing 1
7926979	0.01944806	-0.85024179		
8170009	0.01945979	-0.42042031	FAM127A	family with sequence similarity 127, member A
8135080	0.01946657	-0.37253209	AP1S1	adaptor-related protein complex 1, sigma 1 subunit
8136801	0.01949629	-0.51674739	PRSS3P2///PRSS3///PF	protease, serine, 3 pseudogene 2///protease, serine, 3///protease, serine, 2 (tryp
8046564	0.01951697	-0.32019429	HOXD1	homeobox D1
7893003	0.01952961	-0.58047125		
8038815	0.01955853	-0.36868431	LIM2	lens intrinsic membrane protein 2, 19kDa
7915841	0.01964684	-0.30094815	KNCN	kinocilin
8119712	0.01965196	-0.37958133	SRF	serum response factor (c-fos serum response element-binding transcription fact
8010113	0.0197047	-0.30616097	MGAT5B	mannosyl (alpha-1,6-)-glycoprotein beta-1,6-N-acetyl-glucosaminyltransferase,
7972291	0.01980484	-0.36077884	SOX21	SRY (sex determining region Y)-box 21
8172358	0.01981566	-0.48876675	UXT	ubiquitously-expressed, prefoldin-like chaperone
7898176	0.01986291	-0.3436589	CELA2A	chymotrypsin-like elastase family, member 2A
7979269	0.0198885	-0.33773096	GCH1	GTP cvclohvdrolase 1
8163930	0.01989117	-0.39999006	NDUFA8	NADH dehydrogenase (ubiguinone) 1 alpha subcomplex, 8, 19kDa
7997228	0.01989639	-0.30765144		
7965606	0.01995563	-0.83535876	ΗΔΙ	histidine ammonia-lvase
8109475	0.02002595	-0.41059635	MRPI 22	mitochondrial ribosomal protein I 22
8148104	0.02002000	-0.32845115		
8000200	0.02003972	-0.31126/99		janus kinasa and microtubule interacting protoin 1
7093200	0.02011903	1 270005		CheC section transport regulator homolog 1 (E seli)
7902000	0.02012020	-1.370095		
7909033	0.02014075	-0.35027102	SLAINT	
7892972	0.02014133	-0.45522981	FANADOF	Constitution of the second second second second second
8038109	0.02017002	-0.4/08/3/9	FAM83E	Tarniiy with sequence similarity 83, member E
8180317	0.02021308	-0.65916847	I GIF1	I GFB-induced factor homeobox 1
8037872	0.02022134	-0.38847186	RRC3	BCL2 binding component 3
7914000	0.02023854	-1.01535393	NR0B2	nuclear receptor subfamily 0, group B, member 2
8118804	0.0202519	-0.31237845	PACSIN1	protein kinase C and casein kinase substrate in neurons 1
8047372	0.02028188	-0.47967441	NDUFB3	NADH dehydrogenase (ubiquinone) 1 beta subcomplex, 3, 12kDa
7963970	0.02030429	-0.33986864	PMEL	premelanosome protein
8069532	0 02036576	-0.62754457	HSPA13	heat shock protein 70kDa family, member 13
	0.02000010			
8052669	0.02037158	-0.37371098	SERTAD2	SERTA domain containing 2
8052669 8093074	0.02037158	-0.37371098 -0.36131904	SERTAD2 ZDHHC19	SERTA domain containing 2 zinc finger, DHHC-type containing 19
8052669 8093074 8130009	0.02037158 0.02051172 0.02059968	-0.37371098 -0.36131904 -0.48249909	SERTAD2 ZDHHC19	SERTA domain containing 2 zinc finger, DHHC-type containing 19

8062873	0.02065539	-0.51050214	KCNK15	potassium channel, subfamily K, member 15
7907351	0.02066828	-0.31229037		
7894834	0.02080351	-0.53986824		
7954794	0.02081112	-0.30518875	C12orf40	chromosome 12 open reading frame 40
7969544	0.02084114	-0.36246368	NDFIP2	Nedd4 family interacting protein 2
8136067	0.0210401	-0.38319586	TSPAN33	tetraspanin 33
8134880	0.02107828	-0.37714677	MOSPD3	motile sperm domain containing 3
8076072	0.02110278	-0.41070952	KCNJ4	potassium inwardly-rectifying channel, subfamily J, member 4
7898653	0.02114532	-0.46350768	FAM43B	family with sequence similarity 43, member B
8050790	0.02120541	-0.32776396	DNAJC27	DnaJ (Hsp40) homolog, subfamily C, member 27
7923596	0.02122208	-0.51949514	ETNK2	ethanolamine kinase 2
8168466	0.02123332	-0.34970423	MAGT1	magnesium transporter 1
8175119	0.02123455	-0.30932291		
8110055	0.02124143	-0.31113424	CPEB4	cytoplasmic polyadenylation element binding protein 4
8055404	0.02126076	-0.38070137	UBXN4	LIBX domain protein 4
8052733	0.02134874	-0.33095114	00,000	
7912031	0.02135698	-0 35676742	HES2	hairy and enhancer of split 2 (Drosophila)
7985233	0.02137948	-0 4244739	RASGRE1	Ras protein-specific quanine nucleotide-releasing factor 1
9010793	0.02130015	0.36087111		
8025520	0.02139913	-0.30907111		accomide outbace 1///growth differentiation factor 1
9112726	0.02142094	-0.32391473	CERS1///GDF1	
0112720	0.02142732	-0.30964033		hair and anhancer of anit 1 (Dreagnhile)
0004000	0.02149923	-0.76763537		hairy and enhancer of spirt 1, (Diosophila)
8047217	0.02156229	-0.51903807	COQTOB	coenzyme Q10 nomolog B (S. cerevisiae)
8054831	0.02160232	-0.30893561	EN1	engralied nomeobox 1
7892953	0.0216039	-0.5953891	1000////00	
8088480	0.02161502	-0.40173939	ID2B///ID2	Inhibitor of DNA binding 2B, dominant negative helix-loop-helix protein (pseudog
8012140	0.02162173	-0 42865541	VBX2	V hox binding protein 2
81/5660	0.02166305	-0.3357500/		dynactin 6
7802720	0.02175789	0.55397632	DOTINO	dynactin 6
91292720	0.02173788	-0.33307032		oto voriant 1
0130209	0.02100700	-0.42911011		ets valiant i
8120677	0.02102030	-0.37147413		
0129077	0.02103014	-0.52353047	JUNI	
7925448	0.02184787	-0.36054589	CLIMPOD	akanad multi vasiaulas kartu sestais OD
8081055	0.02190184	-0.3660887		charged multivesicular body protein 2B
8007141	0.02192513	-0.33319703	EIF1	
8120194	0.02208171	-0.30428824	TFAP2B	transcription factor AP-2 beta (activating enhancer binding protein 2 beta)
7986327	0.02215396	-0.39860426		
8128429	0.02218067	-0.42964211	CCNC	cyclin C
7892583	0.0221809	-0.76878003		
7970716	0.02219094	-0.35595354	LNX2	ligand of numb-protein X 2
7944876	0.02224067	-0.43067392	NRGN	neurogranin (protein kinase C substrate, RC3)
8049567	0.02239918	-0.76011616	RAMP1	receptor (G protein-coupled) activity modifying protein 1
7955184	0.02239971	-0.38865949	PRPH	peripherin
8077450	0.02246795	-0.33853098	ARL8B	ADP-ribosylation factor-like 8B
8008540	0.02248531	-0.3707805	FLJ42842	uncharacterized FLJ42842
7967870	0.02249265	-0.50442104	TERF1	telomeric repeat binding factor (NIMA-interacting) 1
7969665	0.0225117	-0.31785763	HS6ST3	heparan sulfate 6-O-sulfotransferase 3
7893778	0.02252415	-0.30331615		
7950671	0.0225325	-0.32449008	GAB2	GRB2-associated binding protein 2
7900146	0.02256338	-0.50669604	ZC3H12A	zinc finger CCCH-type containing 12A
7894004	0.02261292	-0.57197977		
7918869	0.02272015	-0.31672532	NGF	nerve growth factor (beta polypeptide)
7929840	0.02275819	-0.34138901	PAX2	paired box 2
7955719	0.02278816	-0.5853758	HIGD1A	HIG1 hypoxia inducible domain family, member 1A
7987792	0.02280139	-0.30623845	PLA2G4D	phospholipase A2, group IVD (cytosolic)
7991772	0.02283536	-0.37135292	HBQ1	hemoalobin, theta 1
7910387	0.02284822	-0.41487135	RHOU	ras homolog family member U
7928746	0.02288075	-0.42850412		
7896398	0.02289499	-0.56234786		
8016463	0.02305445	-0.33959836	НОХВ6	homeobox B6
80/0509	0.02306453	-0.41366036	ESPNI	asnin-lika
0043030	0.02000400	0.41000000		соритнике

8042251	0.02313198	-0.33084638	OTX1	orthodenticle homeobox 1
8083333	0.02317573	-0.30834586	EIF2A	eukaryotic translation initiation factor 2A, 65kDa
8141303	0.02318773	-0.39082838		
8099172	0.02322339	-0.37595106	CRMP1	collapsin response mediator protein 1
7976496	0.02323646	-0.34662107	SERPINA3	serpin peptidase inhibitor, clade A (alpha-1 antiproteinase, antitrypsin), member
7953697	0.02331522	-0.33217722		
7896324	0.02342616	-0.41711882		
7947221	0.02345421	-0.46365485	LIN7C	lin-7 homolog C (C. elegans)
8003840	0.02346158	-0.42210102	FMC6	ER membrane protein complex subunit 6
7937016	0.02350475	-0.386841	CLRN3	clarin 3
8088889	0.02363846	-0 3373212	0LINIO	our o
8019263	0.02366396	-0 39120489		Rho GDP dissociation inhibitor (GDI) alpha
7894502	0.02371148	-0.53120405	ANIODIA	
7894862	0.02371227	-0.37231410		
7034002 9152512	0.02377635	-0.33013103		tumor pacrosic factor receptor superfamily member 11b
7020256	0.02377033	-0.44393978	INFROFTID	tumor neclosis racior receptor superrainity, member 11b
7920206	0.02376071	-0.52521306		dermaten sulfete enimerose like
8023727	0.02362166	-0.34336667	DSEL	
7903765	0.02388648	-2.80657645	GSTMI	
8091863	0.02395895	-1.44659388	SLITKK3	SLIT and NTKK-like ramily, member 3
7914923	0.02400623	-0.3883317	USCP1	organic solute carrier partner 1
8026163	0.02400683	-0.83125325	IER2	immediate early response 2
7893087	0.02401063	-0.35864595		
8066391	0.02421891	-0.45539776		
8102779	0.02436202	-0.31416575		
8122343	0.02436402	-0.37770049	HECA	headcase homolog (Drosophila)
7980425	0.02437771	-0.31825705	ISM2	isthmin 2
7892697	0.02443476	-0.52723697		
8022488	0.02448474	-0.41883731	ABHD3	abhydrolase domain containing 3
8072488	0.02472195	-0.31448667	DRG1	developmentally regulated GTP binding protein 1
8140140	0.02474001	-0.63393463	CLDN3	claudin 3
8025402	0.02475447	-1.0299977	ANGPTL4	angiopoietin-like 4
8105111	0.02477418	-0.39150814	FBXO4	F-box protein 4
7935270	0.02479891	-0.51852086	BLNK	B-cell linker
8130403	0.02482936	-0.36519141		
8004521	0.02493475	-0.39030054	MPDU1	mannose-P-dolichol utilization defect 1
8170965	0.02496213	-0.31350077	CTAG1A///CTAG2///CT	cancer/testis antigen 1A///cancer/testis antigen 2///cancer/testis antigen 1B
8110872	0.02500576	-0.3790684	IRX2	iroquois homeobox 2
7957611	0.02502437	-0.30675515		
7979033	0.02504748	-0.44091821	SAV1	salvador homolog 1 (Drosophila)
8067820	0.02511803	-0.44436672	IQSEC3	IQ motif and Sec7 domain 3
8133155	0.0251423	-0.37952273	TPST1///TPST1	tvrosvlprotein sulfotransferase 1///tvrosvlprotein sulfotransferase 1
8148265	0.02514553	-0.32541619	RNF139	ring finger protein 139
8135601	0.02514937	-0.42875094	MET	met proto-oncogene (hepatocyte growth factor receptor)
8128716	0.02515417	-0.36886794	CD164	CD164 molecule, sialomucin
8045229	0.025186	-0.42299716	ARHGEF4///ARHGFF4	Rho quanine nucleotide exchange factor (GEF) 4///Rho quanine nucleotide exch
				(GEF) 4
7997556	0.02522968	-0.34025101	DNAAF1///TAF1C	dynein, axonemal, assembly factor 1///TATA box binding protein (TBP)-associat
00/0/	0.007001=1	0.00076	10701	polymerase I, C, 110kDa
8019183	0.02522978	-0.33072592	ACTG1	actin, gamma 1
8147375	0.02526453	-0.46175394	DPY19L4	dpy-19-like 4 (C. elegans)
7892688	0.02537948	-0.56750676		
7975390	0.02538631	-0.41441156	SMOC1	SPARC related modular calcium binding 1
8032312	0.02551534	-0.31898879	ATP8B3	ATPase, aminophospholipid transporter, class I, type 8B, member 3
8068593	0.02552488	-0.71143207	ETS2	v-ets erythroblastosis virus E26 oncogene homolog 2 (avian)
8022418	0.02553354	-0.45049931		
8137485	0.02555227	-0.33073436	DPP6	dipeptidyl-peptidase 6
8169210	0.02568072	-0.56449138	RIPPLY1///CLDN2	ripply1 homolog (zebrafish)///claudin 2
8150908	0.02570623	-0.39409707	IMPAD1	inositol monophosphatase domain containing 1
8102594	0.02570979	-0.37260471	TNIP3	TNFAIP3 interacting protein 3
8056977	0.02572337	-0.3593071	NFE2L2	nuclear factor (erythroid-derived 2)-like 2
8066590	0.02577605	-0.36891177	TNNC2	troponin C type 2 (fast)
8033162	0.02578024	-0.30152477	KHSRP	KH-type splicing regulatory protein
8079163	0.02580835	-0.58638954		· · · ·
			1	

8100306	0.02590369	-0.37449119		
7895217	0.02592048	-0.4714582		
7896310	0.02593443	-0.30808722		
7894704	0.02594183	-0.38018161		
8083415	0.02598718	-0.44414187	AADAC	arvlacetamide deacetvlase
8061944	0.02600145	-0.34539848	ACTI 10	actin-like 10
7893173	0.02601082	-0.69304519	101210	
7001808	0.02606118	-0.03304313		NHL repeat containing 4
9144279	0.0261002	0.4740334		1 opulativesrel 2 phosphoto O coultransformer 5
0144370	0.0201092	-0.43346736		
8079170	0.02614056	-0.33176241		
0100040	0.02015140	-0.95555334	NR4A3	nuclear receptor subramily 4, group A, member 3
8099506	0.02625425	-0.3595168		transmembrane anterior posterior transformation 1
8146717	0.02626105	-0.32853884	SGK3	serum/glucocorticoid regulated kinase family, member 3
8149720	0.02639784	-0.53197671	EGR3	early growth response 3
7893619	0.0264725	-0.50591131		
8036133	0.02651955	-0.32277986	UPK1A	uroplakin 1A
8062539	0.02652947	-0.32068848	SLC32A1	solute carrier family 32 (GABA vesicular transporter), member 1
7914354	0.02656147	-0.34405917	PEF1	penta-EF-hand domain containing 1
8042962	0.02658871	-0.42156346	MRPL19	mitochondrial ribosomal protein L19
7903162	0.02660328	-0.31385927	TMEM56	transmembrane protein 56
7895973	0.02661055	-0.34212364		
8073007	0.02671792	-0.57984919	MAFF	v-maf musculoaponeurotic fibrosarcoma oncogene homolog F (avian)
8111788	0.02673459	-0.38484986	TTC33	tetratricopeptide repeat domain 33
8044301	0.02674857	-0.30485791	SOWAHC	sosondowah ankyrin repeat domain family member C
8131803	0.0268214	-0.89782804	IL6	interleukin 6 (interferon, beta 2)
7896255	0.0268285	-0.49141501		
7913146	0.02684534	-0.30739874	AKR7A3	aldo-keto reductase family 7, member A3 (aflatoxin aldehyde reductase)
8064779	0.0268465	-0.34906901	ADRA1D	adrenoceptor alpha 1D
7911376	0.02686542	-0.31042638	HES4	hairy and enhancer of split 4 (Drosophila)
7941172	0.02693658	-0.36190574	SPDYC	speedv/RINGO cell cvcle regulator family member C
8138527	0.0269443	-0.34457007	STEAP1B	STEAP family member 1B
8172030	0.02694978	-0.30162403		,
7969479	0.02704771	-0.31087773	BTF3P11	basic transcription factor 3 pseudogene 11
7975626	0.02706253	-0.45496525	ELMSAN1	ELM2 and Mvb/SANT-like domain containing 1
7997712	0.02712303	-0.72653936	IRF8	interferon regulatory factor 8
7976858	0.02733543	-0.35806974	DIO3	deiodinase, iodothyronine, type III
8052698	0.02736741	-0.3432308	C1D	C1D nuclear receptor corepressor
7963760	0.02745965	-0.34943116	NFF2	nuclear factor (erythroid-derived 2) 45kDa
8142697	0.0275065	-0 38058452	POT1	protection of telomeres 1
789/355	0.02752803	-0.47717503	1011	
7094333	0.02755705	0.30120457		
7030200	0.02100100	-0.33120437	MXI1	MAX interactor 1. dimerization protein
9177046	0.02770021	0.55266202		אות אווופומטוטר ז, שווופוצמווטוו אוטופוו
7907024	0.02779921	-0.000000000000000000000000000000000000	CARRO	anno aminahuturia aaid (CADA) A recentar dalta
109/034	0.02700500	-0.31023434		yanima-animobulync aciu (GADA) A receptor, delta
01/1023	0.02789586	-0.34248416		aponpoprotern O
0032834	0.02793904	-0.00454653		teucine-rich alpha-2-glycoprotein 1
8065353	0.02795063	-0.46532419	IHRD	nindomoduin
/896690	0.0280276	-0.50665589		
8001798	0.02805468	-0.31075312	1 0 0 0 1 C 1 = 1 11 C ====	
7913850	0.02810734	-0.33194813	LOC646471///MTFR1L	uncharacterized LOC646471///mitochondrial fission regulator 1-like
8002999	0.02810917	-0.31232508	GCSHP3///LOC729080	glycine cleavage system protein H (aminomethyl carrier) pseudogene 3///glycine
				system protein H (aminometnyi carrier) pseudogene///glycine cleavage system p
7941269	0 02816762	-0.39682246		
8123728	0.0281749	-0.35133147	LYRM4	LYR motif containing 4
8078/35	0.02818281	-0.32867305	TRIM71	trinartite motif containing 71 F3 ubiquitin protain ligase
7065040	0.02010201	-0.02007303		njarke moli containing / 1, Lo ubiquilli protein ligase
7017700	0.02013430	0.3033430		formity with sequence similarity 60, member A
1311120 9010007	0.02020300	0.0042000		chromosomo 17 opon roading from 00
0010237	0.02831182			chromosome 17 open reading frame 99
00238/1	0.02831187	-0.31051338		zine binding alconol denydrogenase domain containing 2
/895334	0.02831804	-0.406/1984	DAVA	natural have d
8061357	0.02832728	-0.41/37689	PAX1	paired box 1

7955441	0.02841787	-0.34617998	METTL7A	methyltransferase like 7A	
7922051	0.02846687	-0.36141561	CREG1	cellular repressor of E1A-stimulated genes 1	
7894304	0.02854155	-0.43810697			
8016540	0.02857595	-0.38617005	PHOSPHO1	phosphatase, orphan 1	
7901802	0.02859746	-0.31103064	MGC34796	sepiapterin reductase (7,8-dihydrobiopterin:NADP+ oxidoreductase) pseudogen	
8121144	0.02862334	-0.53181855	MANEA	mannosidase, endo-alpha	
8168578	0.0286449	-0.32061114	UBE2DNL	ubiquitin-conjugating enzyme E2D N-terminal like (pseudogene)	
7968236	0.02879759	-0.32860821	RASL11A	RAS-like, family 11, member A	
8076677	0.02881079	-0.32440831	PHF21B	PHD finger protein 21B	
7915329	0.02889301	-0.35165665			
7980523	0.02894642	-0.36038494	GTF2A1	general transcription factor IIA, 1, 19/37kDa	
8010036	0.02915004	-0.31136209	ZACN///EXOC7///EXOC	zinc activated ligand-gated ion channel///exocyst complex component 7///exocyst component 7	
8151125	0.02920851	-0.38130249	TCF24	transcription factor 24	
7936826	0.0293103	-0.49470972	IKZF5	IKAROS family zinc finger 5 (Pegasus)	
8141066	0.0293129	-0.34087125	PON3	paraoxonase 3	
8083324	0.02931723	-0.44598425	TSC22D2	TSC22 domain family, member 2	
7984174	0.02934602	-0.36266321	SNX22///PPIB	sorting nexin 22///peptidylprolyl isomerase B (cyclophilin B)	
8106336	0.02935051	-0.32301615	SV2C	synaptic vesicle glycoprotein 2C	
8065403	0.02937605	-0.34332977	CST3	cystatin C	
7935692	0.02946431	-0.34669087	ERLIN1	ER lipid raft associated 1	
8166569	0.02947284	-0.38794584			
8075118	0.02950802	-0.37285849	CRYBB1	crystallin, beta B1	
7893082	0.02963252	-0.72359235			
7893928	0.02963398	-0.44527055			
8099073	0.02966075	-0.36510382	LRPAP1	low density lipoprotein receptor-related protein associated protein 1	
8112592	0.02969395	-0.30322489	FOXD1	forkhead box D1	
7914974	0.02974117	-0.31862742	GRIK3	glutamate receptor, ionotropic, kainate 3	
7990090	0.02983301	-0.30991691			
8024934	0.02984721	-0.30359383	ZNRF4	zinc and ring finger 4	
8037003	0.02989863	-0.34628531			
8086729	0.02990948	-0.34223589	KIF9	kinesin family member 9	
7914180	0.03003791	-0.31162767	SPCS2	signal peptidase complex subunit 2 homolog (S. cerevisiae)	
7914180 8175288	0.03003791	-0.31162767 -0.49348638	SPCS2 MOSPD1	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1	
7914180 8175288 8142307	0.03003791 0.03006719 0.03007847	-0.31162767 -0.49348638 -0.46495251	SPCS2 MOSPD1 PNPLA8	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8	
7914180 8175288 8142307 7974207	0.03003791 0.03006719 0.03007847 0.03022414	-0.31162767 -0.49348638 -0.46495251 -0.33246294	SPCS2 MOSPD1 PNPLA8 MGAT2	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1.6-)-glycoprotein beta-1.2-N-acetylglucosaminyltransferase	
7914180 8175288 8142307 7974207 8109505	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3	
7914180 8175288 8142307 7974207 8109505 7956613	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31	
7914180 8175288 8142307 7974207 8109505 7956613 8114778	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine c	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.03044569	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.03044569 0.03045036	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine c protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.03044569 0.03045036 0.03052556	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine c protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526 8037205	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.03044569 0.03044569 0.03045036 0.03052556 0.03060711	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035 -0.37469774	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2 CEACAM1	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary dycoprotein)	
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7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526 8037205 8085116 8041000 8157246 8162848 7893259 7975268 7969414 7896667 8166202 7910001 7974229 7945781	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.0304569 0.03044569 0.03044569 0.03044569 0.03044569 0.0304556 0.03060711 0.03069795 0.0308347 0.03093764 0.03102888 0.03106309 0.03108553 0.03118009 0.03119811 0.03148209 0.0315067 0.0315067	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035 -0.37469774 -0.4270489 -0.34092652 -0.30004733 -0.46982795 -0.45965939 -0.67223504 -0.75072228 -0.30314426 -0.31081949 -0.34819306 -0.32930146 -0.32735019 -0.4275242	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2 CEACAM1 EDEM1 GPN1 KIAA1958 VTI1B///ARG2 VTI1B///ARG2 KLF5 GRPR DEGS1 KLHDC2 PHLDA2 CNIDV2	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein) ER degradation enhancer, mannosidase alpha-like 1 GPN-loop GTPase 1 KIAA1958 vesicle transport through interaction with t-SNAREs 1B///arginase 2 Kruppel-like factor 5 (intestinal) gastrin-releasing peptide receptor delta(4)-desaturase, sphingolipid 1 kelch domain containing 2 pleckstrin homology-like domain, family A, member 2 ensony (2 homolog, (rabrafish)	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526 8037205 8085116 8041000 8157246 8162848 7893259 7975268 7969414 7896667 8166202 7910001 7974229 7945781 8119609 8454422	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.0304569 0.03044569 0.03044569 0.03044569 0.03045036 0.03052556 0.03060711 0.03069795 0.03069795 0.0308347 0.03093764 0.03102888 0.03106309 0.03108553 0.03118009 0.03118019 0.03148209 0.0315067 0.03162741 0.03162755	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035 -0.37469774 -0.4270489 -0.34092652 -0.30004733 -0.46982795 -0.45965939 -0.67223504 -0.75072228 -0.30314426 -0.31081949 -0.34819306 -0.32735019 -0.43975818	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2 CEACAM1 EDEM1 GPN1 KIAA1958 VTI1B///ARG2 KLF5 GRPR DEGS1 KLHDC2 PHLDA2 CNPY3 DCL4	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein) ER degradation enhancer, mannosidase alpha-like 1 GPN-loop GTPase 1 KIAA1958 vesicle transport through interaction with t-SNAREs 1B///arginase 2 Kruppel-like factor 5 (intestinal) gastrin-releasing peptide receptor delta(4)-desaturase, sphingolipid 1 kelch domain containing 2 pleckstrin homology-like domain, family A, member 2 canopy 3 homolog (zebrafish)	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526 8037205 8085116 804200 8157246 8162848 7893259 7975268 7969414 7896667 8166202 7910001 7974229 7945781 8119609 8154163 7004555	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.0304569 0.03044569 0.03044569 0.03044569 0.03045036 0.03052556 0.03060711 0.03069795 0.0308347 0.03093764 0.03108553 0.03108553 0.03119811 0.03140884 0.03146211 0.03146211 0.03165353 0.0316577 0.03165741 0.03165353	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035 -0.37469774 -0.4270489 -0.34092652 -0.30004733 -0.46982795 -0.45965939 -0.67223504 -0.75072228 -0.30314426 -0.31081949 -0.34819306 -0.32735019 -0.43975818 -0.36892988	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2 CEACAM1 EDEM1 GPN1 KIAA1958 VTI1B///ARG2 VTI1B///ARG2 KLF5 GRPR DEGS1 KLHDC2 PHLDA2 CNPY3 RCL1	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine of protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein) ER degradation enhancer, mannosidase alpha-like 1 GPN-loop GTPase 1 KIAA1958 vesicle transport through interaction with t-SNAREs 1B///arginase 2 Kruppel-like factor 5 (intestinal) gastrin-releasing peptide receptor delta(4)-desaturase, sphingolipid 1 kelch domain containing 2 pleckstrin homology-like domain, family A, member 2 canopy 3 homolog (zebrafish) RNA terminal phosphate cyclase-like 1	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526 8037205 8085116 8041000 8157246 8162848 7893259 7975268 7969414 7896667 8166202 7910001 7974229 7945781 8119609 8154163 7924553	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.03044569 0.03044569 0.03044569 0.03045036 0.03052556 0.03060711 0.03069795 0.0308347 0.030093764 0.03102888 0.03106309 0.03108553 0.03118009 0.03148209 0.03162741 0.03165353 0.03173748 0.03173748	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035 -0.37469774 -0.4270489 -0.34092652 -0.30004733 -0.46982795 -0.45965939 -0.67223504 -0.75072228 -0.30314426 -0.31081949 -0.34819306 -0.32930146 -0.32735019 -0.43975818 -0.36892988 -0.37378922	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2 CEACAM1 EDEM1 GPN1 KIAA1958 VTI1B///ARG2 KLF5 GRPR DEGS1 KLHDC2 PHLDA2 CNPY3 RCL1 LUEM0	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine c protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein) ER degradation enhancer, mannosidase alpha-like 1 GPN-loop GTPase 1 KIAA1958 vesicle transport through interaction with t-SNAREs 1B///arginase 2 Kruppel-like factor 5 (intestinal) gastrin-releasing peptide receptor delta(4)-desaturase, sphingolipid 1 kelch domain containing 2 pleckstrin homology-like domain, family A, member 2 canopy 3 homolog (zebrafish) RNA terminal phosphate cyclase-like 1	
7914180 8175288 8142307 7974207 8109505 7956613 8114778 8015376 8060594 7974531 8042283 7924526 8037205 8085116 8041000 8157246 8162848 7893259 7975268 7969414 7896667 8166202 7910001 7974229 7945781 8119609 8154163 7924553 8148559	0.03003791 0.03006719 0.03007847 0.03022414 0.03025007 0.03028358 0.03032904 0.03035113 0.03039813 0.03044569 0.03044569 0.03045036 0.03052556 0.03060711 0.03069795 0.0308347 0.03093764 0.03108553 0.03108553 0.03108553 0.03119811 0.03146211 0.03146211 0.03165353 0.0315067 0.03162741 0.03165353 0.03173748 0.03178579 0.03178579	-0.31162767 -0.49348638 -0.46495251 -0.33246294 -0.3168818 -0.32320653 -0.31923659 -0.35383165 -0.31677192 -0.40690066 -0.4313904 -0.5730035 -0.37469774 -0.4270489 -0.34092652 -0.30004733 -0.46982795 -0.45965939 -0.67223504 -0.75072228 -0.30314426 -0.31081949 -0.34819306 -0.32930146 -0.32735019 -0.43975818 -0.36892988 -0.37378922 -0.3286659 -0.228659	SPCS2 MOSPD1 PNPLA8 MGAT2 PPP1R2P3 TSPAN31 LOC729080///GCSH KRT16 GNRH2 RPL13AP3 LGALSL TP53BP2 CEACAM1 EDEM1 GPN1 KIAA1958 VTI1B///ARG2 KLF5 GRPR DEGS1 KLHDC2 PHLDA2 CNPY3 RCL1 THEM6 ADD	signal peptidase complex subunit 2 homolog (S. cerevisiae) motile sperm domain containing 1 patatin-like phospholipase domain containing 8 mannosyl (alpha-1,6-)-glycoprotein beta-1,2-N-acetylglucosaminyltransferase protein phosphatase 1, regulatory (inhibitor) subunit 2 pseudogene 3 tetraspanin 31 glycine cleavage system protein H (aminomethyl carrier) pseudogene///glycine c protein H (aminomethyl carrier) keratin 16 gonadotropin-releasing hormone 2 ribosomal protein L13a pseudogene 3 lectin, galactoside-binding-like tumor protein p53 binding protein, 2 carcinoembryonic antigen-related cell adhesion molecule 1 (biliary glycoprotein) ER degradation enhancer, mannosidase alpha-like 1 GPN-loop GTPase 1 KIAA1958 vesicle transport through interaction with t-SNAREs 1B///arginase 2 Kruppel-like factor 5 (intestinal) gastrin-releasing peptide receptor delta(4)-desaturase, sphingolipid 1 kelch domain containing 2 pleckstrin homology-like domain, family A, member 2 canopy 3 homolog (zebrafish) RNA terminal phosphate cyclase-like 1	

7997569	0.03183536	-0.3593739	ADAD2	adenosine deaminase domain containing 2
7989365	0.03184464	-0.35315308	RORA	RAR-related orphan receptor A
8096905	0.03194318	-0.3484977	AP1AR	adaptor-related protein complex 1 associated regulatory protein
7894138	0.0319447	-0.41249327		
8112072	0.03205433	-0.32973633	CCNO	cyclin O
7919582	0.03215064	-0.45153478		
8062461	0.0321613	-0.4378055	LBP	lipopolysaccharide binding protein
7909127	0.03217148	-0.30164741	MFSD4	maior facilitator superfamily domain containing 4
8004241	0.03218767	-0.30138601	RNASEK///C17orf49	ribonuclease, RNase K///chromosome 17 open reading frame 49
7925950	0.03226732	-0.44792722	UCN3	urocortin 3
7905929	0.03228423	-0.46237758	FFNA1	ephrin-A1
8131406	0.03229614	-0.43798212	RAC1	ras-related C3 botulinum toxin substrate 1 (rbo family, small GTP binding protein
8029701	0.03231542	-0.56255539	PPM1N	protain phosphatase Mr2+/Mr2+ dependent 1N (putative)
7950032	0.03231706	-0.32374714	FGF4	fibroblast growth factor 4
7895685	0.03232665	-0.38863908		
7001883	0.03232003	-0.30073131		angiopoletin-like 3
7901003	0.03235727	1 10024620		
9157120	0.032430	-1.19924029	3011	
0107109	0.03247020	-0.39621663		
01/4091	0.03251936	-0.34442649		
8015037	0.0326976	-0.4437154	007142	hutathiana C transforman mu 2 (husin)
7918379	0.03271243	-1.06526659		giutatnione S-transferase mu 3 (brain)
7912670	0.03271953	-0.41278526	UQCRHL///UQCRH	ubiquinoi-cytochrome c reductase ninge protein-like///ubiquinoi-cytochrome c re-
8089013	0.03272423	-0.44129858		
8129254	0.03277701	-0.31788096	MAN1A1	mannosidase, alpha, class 1A, member 1
8161648	0.03279122	-0.58184507	KI F9	Kruppel-like factor 9
7990729	0.03282401	-0 32144785	CHRNB4	cholineraic receptor nicotinic beta 4 (neuronal)
8128001	0.03287032	-0.36487137	CGA	alvoorotain bormones, alpha polypentide
7007733	0.03280306	-0.30153026	EOXC2	forkhead box C2 (MEH-1, mesenchyme forkhead 1)
8012500	0.03203330	0.30154468	C17orf103	chromosomo 17 opon roading framo 103
7904027	0.03292713	-0.30134408	017011103	
1094921	0.03314063	-0.49501701		alaudia 22
8144505	0.03314709	-0.31133007		ciaudin 23
7994804	0.03315019	-0.30419955		ADD site and the set of the set o
7961507	0.0331577	-0.41364629	AR14	ADP-ribosyltransierase 4 (Dombrock blood group)
8076561	0.03322706	-0.37209892	KIAA1654	KIAA1654 protein
8066031	0.033325	-0.30769332	SCAND1	SCAN domain containing 1
8048699	0.03339672	-0.37034553	TIDINI	
7989915	0.03350831	-0.35760344		TIMELESS interacting protein
7979179	0.03356123	-0.49963529	ERO1L	ERO1-like (S. cerevisiae)
8076128	0.0336813	-0.40784078	JOSD1	Josephin domain containing 1
8070169	0.03371285	-0.32026315		
8096538	0.03381924	-0.39575948	RAP1GDS1	RAP1, GTP-GDP dissociation stimulator 1
8129637	0.03384934	-0.71846815	VNN2	vanin 2
8034578	0.03392588	-0.44430408	KLF1	Kruppel-like factor 1 (erythroid)
8172573	0.03396907	-0.42796571	SYP	synaptophysin
8043718	0.0340294	-0.35804576	COX5B	cytochrome c oxidase subunit Vb
7895873	0.03416901	-0.54949333		
8154848	0.03420829	-0.38688874	PRSS3///PRSS2	protease, serine, 3///protease, serine, 2 (trypsin 2)
8038782	0.03425367	-0.34151532	CTU1	cytosolic thiouridylase subunit 1
7892808	0.03429172	-0.35178188		
8046086	0.03432726	-0.47280196	CERS6	ceramide synthase 6
7930311	0.03451604	-0.38494613	GSTO2	glutathione S-transferase omega 2
7956668	0.03452285	-0.35508016		
7943760	0.03458118	-0.32568226	SIK2	salt-inducible kinase 2
8139889	0.03473559	-0.57016099		
8066513	0.03475014	-0.50765406	SDC4	syndecan 4
8071809	0.03475814	-1.11041942	GSTT2B///GSTT2	glutathione S-transferase theta 2B (gene/pseudogene)///glutathione S-transferase
8156058	0.03477317	-0.3385584		
7932794	0.03479717	-0.43998284		
8070863	0.03481119	-0.32447879	SSR4P1	signal sequence receptor, delta pseudogene 1
7926674	0.03486934	-0.39434239	PTF1A	pancreas specific transcription factor, 1a
7898673	0.03489448	-0.38741235		
		====		

7992678	0.03498186	-0.43720439	LOC652276	potassium channel tetramerisation domain containing 5 pseudogene
7894599	0.03506526	-0.8500798		
8100994	0.03521137	-0.89961072	CXCL2	chemokine (C-X-C motif) ligand 2
7896079	0.03523457	-0.78866393		
7945101	0.03531058	-0.33274163	DCPS	decapping enzyme, scavenger
8105302	0.03539902	-1.07895136	FST	follistatin
8039892	0.03575151	-0.31856104	KIR2DS5///KIR2DS3	killer cell immunoglobulin-like receptor, two domains, short cytoplasmic tail, 5///k immunoglobulin-like receptor, two domains, short cytoplasmic tail, 3
7907859	0.03583894	-0.36042964		
8170633	0.03585005	-0.41274458		
8175773	0.03585005	-0.41274458		
8084963	0.03587451	-0.5106764	PAK2	p21 protein (Cdc42/Rac)-activated kinase 2
8000467	0.03590794	-0.33572319	GSG1L	GSG1-like
7893500	0.03604075	-0.36131196		
8040547	0.03609127	-0.36165176		
8011990	0.03610252	-0.3392767	TEKT1	tektin 1
8034565	0.03613846	-0.35546524	DNASE2	deoxyribonuclease II, lysosomal
7924758	0.03615374	-0.68660195		
8143788	0.0362049	-0.31173326		
7941122	0.03622614	-0.42196295	SAC3D1	SAC3 domain containing 1
7968734	0.03622724	-0.430126	SLC25A15	solute carrier family 25 (mitochondrial carrier; ornithine transporter) member 15
8166455	0.03628888	-0.37811587	PRDX4	peroxiredoxin 4
8135392	0.03634644	-0.31292591	HBP1	HMG-box transcription factor 1
8018864	0.03637402	-0.97614025	SOCS3	suppressor of cytokine signaling 3
8130715	0.03637743	-0.31640538	PRR18	proline rich 18
8107100	0.03637959	-0.52635217	RGMB	RGM domain family, member B
7919600	0.0364701	-1.18776629		
8180277	0.03649965	-0.41422308	PCDHA3	protocadherin alpha 3
7970563	0.03653589	-0.3664252		F
8107671	0.03656598	-0.49789462		
8111216	0.03662959	-0.31002799		
7999360	0.03669984	-0.35232371	RPI 21P28///RPI 21	ribosomal protein I 21 pseudogene 28///ribosomal protein I 21
8005097	0.03674099	-0.41900767	HS3ST3B1	henaran sulfate (ducosamine) 3-O-sulfotransferase 3B1
8024728	0.03680636	-0.4029797	NMRK2	nicotinamide riboside kinase 2
8108066	0.03683949	-0.30291868	LIBE2B	ubiquitin-conjugating enzyme E2B
8166049	0.03687075	-0.31750506	PRPS2	nhosnhorihosyl nyronhosnhate synthetase 2
8040516	0.0369157	-0 5051371	MESD2B	major facilitator superfamily domain containing 2B
7974387	0.03694516	-0 31008798	STYX	serine/threonine/throsine interacting protein
8161190	0.03703269	-0.49644051	UTIX	
8077441	0.03709841	-0.49044031		hasic balix-loop-balix family, member e40
80/0187	0.03703041	0.47519107		EE band domain family, member D1
7896070	0.0372107	-0.33076845		
8060676	0.03722757	0 85870656		ADAM matallapoptidase with thrombospondin type 1 matif 1
8000201	0.037/3618	-0.03079030	ALGI	ALC1 chitobiosyldinbosphodolichol beta-mannosyltransferase-like
7803/2/	0.03743010	-0.41000200		הבסיו, טוונטאוטאוטאוטאוטטאוטעטוטוטו שבומ-ווומוווטאונומוואובומשביווגל
8018705	0.03747201	-0.33342230		
7806000	0.03747207	0.01007020		
7070710	0.03740422	-0.0209/300	EBY022	E hov protoin 22
1310140	0.03752102	-0.4000/441	TMEM200	transmembrane protain 200
0142912	0.03752103	-0.30271770		
0042100 2022525	0.03750364	-0.01907	AUTUZ SLC2042	aum, yanima 2, Sinouti musule, entenu soluto corrior family 20 (zing transporter), member 2
80002020	0.03760603	-0.30073297		Soule camer raminy 59 (zinc transporter), member 5
0000323	0.03761989	-0.310438/0		North denydrogenase (dbiquinone) 1, alpha/beta subcomplex, 1, 8kDa
8143040	0.03762051	-0.33467517	3LU33D4	Solute carrier family 35, member B4
0042115	0.03700376	-0.31040491		
79/4253	0.03766438	-0.30503/0/		
7090544	0.03/6/489	-0.42552135		colled coll demain containing 470///aldebude debude services 0.(coll
7980098	0.0377386	-0.3229128	CCDC176///ALDH6A1//	colled-coll domain containing 176///aldenyde dehydrogenase 6 family, member / dehydrogenase 6 family, member A1
8151056	0.03776473	-0.44895028	СҮР/В1	cytochrome P450, tamily /, subfamily B, polypeptide 1
8162610	0.03790572	-0.31454809	CDC14B	cell division cycle 14B
8046680	0.03792176	-0.37746179	PLEKHA3	pleckstrin homology domain containing, family A (phosphoinositide binding spec
8148553	0.03792385	-0.33180889	LY6K	lymphocyte antigen 6 complex, locus K
7893640	0.03796471	-0.47023799		

7895180	0.0380123	-0.39490714		
7915084	0.03802175	-0.41401087	YRDC	yrdC domain containing (E. coli)
8131539	0.03805652	-0.42472604	TMEM106B	transmembrane protein 106B
7965510	0.03818661	-0.37599997	TMCC3	transmembrane and coiled-coil domain family 3
7892839	0.03828469	-0.52130725		
8086462	0.03828507	-0.34427825	GTDC2	glycosyltransferase-like domain containing 2
8146930	0.03831349	-0.37505013	TMEM70	transmembrane protein 70
7981309	0.03833545	-0.37944993	BEGAIN	brain-enriched guanylate kinase-associated
8157818	0.03840628	-0.34990148	WDR38	WD repeat domain 38
7896230	0.03847837	-0.32376766		
7998910	0.03849053	-0.3726705	CCDC64B	coiled-coil domain containing 64B
8088745	0.03856495	-0.40317011	FRMD4B	FERM domain containing 4B
8117045	0.0385961	-0.42829137	RBM24	RNA binding motif protein 24
7922717	0.03870875	-0 40204553	RGS16	regulator of G-protein signaling 16
8036557	0.0387583	-0 44289261		
8006569	0.03883207	-0.32168146	C17orf50	chromosome 17 open reading frame 50
7894519	0.0389427	-0.45007921	0110100	
7965508	0.03895378	-0.43266683		
9167990	0.03003070	0.30822552	PAGES	P antigan family, member 5 (prestate associated)
7902661	0.03902030	-0.30022332	FAGLS	F antigen family, member 5 (prostate associated)
9126215	0.0390339	-0.34042073		
0130213	0.03911033	-0.33367642		protoin interacting with evolin A1
8013747	0.03911946	-0.30577509		protein interacting with cyclin A1
8151296	0.03918597	-0.44884649		lactamase, beta 2
8146115	0.03926141	-0.70187996	C80ff4	chromosome 8 open reading frame 4
7936242	0.03927188	-0.32557283		inositol 1,4,5-trisphosphate receptor interacting protein
7917088	0.03939231	-0.47481236	PIGK	phosphatidylinositol glycan anchor biosynthesis, class K
7969651	0.03940047	-0.42909617	DNAJC3	DnaJ (Hsp40) homolog, subfamily C, member 3
7957467	0.03940386	-0.33457218	C12orf29///CEP290	chromosome 12 open reading frame 29///centrosomal protein 290kDa
7939465	0.0394497	-0.32307716	HSD17B12	hydroxysteroid (17-beta) dehydrogenase 12
8125316	0.03948778	-0.31327916	FKBPL	FK506 binding protein like
8043320	0.03967198	-0.42123852	LOC400965	uncharacterized LOC400965
7902074	0.03978258	-0.53527499	LEPROT///LEPR	leptin receptor overlapping transcript///leptin receptor
8139879	0.04009814	-0.40630081	SKP1	S-phase kinase-associated protein 1
8043022	0.0402328	-0.3720733		
7898693	0.04023794	-0.57118973	ALPL	alkaline phosphatase, liver/bone/kidney
8072113	0.04024057	-0.3599734	SRRD	SRR1 domain containing
7892682	0.04028159	-0.61767811		
8132897	0.04031903	-0.38316288	LANCL2	LanC lantibiotic synthetase component C-like 2 (bacterial)
7895149	0.04033131	-0.35104922		
8021707	0.04036137	-0.40527748	SOCS6	suppressor of cytokine signaling 6
7960253	0.04036618	-0.40279901	NINJ2	ninjurin 2
8180373	0.04044063	-0.30530173	SUMO1	small ubiguitin-like modifier 1
8131881	0.04050256	-0.51546753	FAM221A	family with sequence similarity 221, member A
7909888	0.04055082	-0.39481768		
7895876	0.04064224	-0.51776048		
7894830	0.04078726	-0.38072844		
8133704	0.04086501	-0.31752423	SRRM3///SRRM3	serine/arginine repetitive matrix 3///serine/arginine repetitive matrix 3
8143949	0.04092381	-0.31269606	CRYGN	crystallin, gamma N
8026292	0.04102516	-0.33918384	EFE1D	eukaryotic translation elongation factor 1 delta (quanine nucleotide exchange pr
8051215	0.04102310	-0.33910304	BRE-AS1///RBKS	BRE antisense RNA 1///ribokinase
7802603	0.04100002	0.57770842	DICE-AG I/// ICDICG	
7092003	0.04119407	-0.37770042		
7693641	0.04123141	-0.41225562		untral antarian hamaahay 0
8042532	0.04131916	-0.31021714		
0090891	0.04135466	-0.33407575		EPH receptor BT
81145/2	0.04145722	-0.69304903	HBEGE	neparin-binding EGF-like growth factor
/930304	0.04152995	-0.44213784	GSTO1	giutathione S-transferase omega 1
7894542	0.04163326	-0.45839945		
7978628	0.04169152	-0.303697	PPP2R3C	protein phosphatase 2, regulatory subunit B", gamma
8097461	0.04169451	-0.94018506	CCRN4L///CCRN4L	CCR4 carbon catabolite repression 4-like (S. cerevisiae)///CCR4 carbon catabol
7040404	0.04474770	0.20020025		IIKe (5. cerevisiae)
1910494	0.041/1//8	-0.38933035		
8096733	0.041/5978	-0.38660729	SGMS2	spningomyelin synthase 2

7933872	0.0418679	-0.67145235	EGR2	early growth response 2
8000998	0.04187791	-0.3586624	VKORC1	vitamin K epoxide reductase complex, subunit 1
8105852	0.0419086	-0.32586931	MRPS36	mitochondrial ribosomal protein S36
7982574	0.04192064	-0.35892474	FAM98B	family with sequence similarity 98, member B
7978846	0.04198855	-0.41409048	POLE2	polymerase (DNA directed), epsilon 2, accessory subunit
7984405	0.04211497	-0.46112505	C15orf61	chromosome 15 open reading frame 61
8156761	0.04219894	-0.3833524	NANS	N-acetylneuraminic acid synthase
8038899	0.04239774	-0.51731989	FPR1	formyl peptide receptor 1
8065327	0.04247344	-0.43658117	NKX2-4	NK2 homeobox 4
7896060	0.04249677	-0.40557897		
8051949	0.04278211	-0.4063265	SIX2	SIX homeobox 2
7919564	0.04299338	-0.53504869	•=	
7984103	0.04300036	-0 33968794	LACTB	lactamase beta
8008980	0.04302428	-0 41815642	C17orf82	chromosome 17 open reading frame 82
8067111	0.04316854	-0 34170788	0 IT ONOL	
7894165	0.04336137	-0.43868054		
8113369	0.04337134	-0.85935617	SI CO4C1	solute carrier organic anion transporter family, member 4C1
8037179	0.04340699	-0.36773145	CNEN	
7964660	0.04343101	-1.04348582		arginine vasopressin recentor 1A
7904000	0.04345025	0.20608576		
9067912	0.04343923	-0.30090370	C20orf201	chromosomo 20 opon reading fromo 201
815/125	0.04303190	-0.00004009		solute carrier family 1 (neuropal/anithalia) high affinity dutemate transporter, and
0104100	0.04307493	-0.40700071	SLUTAT	member 1
8160531	0.04367916	-0.46596527	C9orf72	chromosome 9 open reading frame 72
7896581	0.04376155	-0.60576349		
8119161	0.04379688	-0.3655529	PIM1	pim-1 oncogene
7898594	0.04380584	-0.31414433	HTR6	5-hydroxytryptamine (serotonin) receptor 6. G protein-coupled
7953032	0.04385456	-0.31093702	LRTM2	leucine-rich repeats and transmembrane domains 2
8054154	0.0439285	-0.31709247	KIAA1211I	KIAA1211-like
8039269	0.04396328	-0.33332001	LENG8///LENG9	leukocyte receptor cluster (LRC) member 8///leukocyte receptor cluster (LRC) m
7930208	0.0439735	-0.32327182	INA	internexin neuronal intermediate filament protein, alpha
8023246	0.04427122	-0.44347621	C18orf32	chromosome 18 open reading frame 32
7962884	0.04428387	-0.91955532	RND1	Rho family GTPase 1
7935011	0.04430029	-0.37042566	CPEB3	cytoplasmic polyadenylation element binding protein 3
8057045	0.04431619	-0.35951489	FKBP7	EK506 binding protein 7
7897424	0.0443265	-0.31869502		
7975284	0.04434487	-0.37654479	RDH12	retinol dehydrogenase 12 (all-trans/9-cis/11-cis)
7892526	0.04444113	-0.54199907	1101112	
8103166	0.04454597	-0 36943995	SH3D19	SH3 domain containing 19
8162833	0.04468221	-0.33186974	ERP44	endonlasmic reticulum protein 44
700/061	0.04472426	-0.30552266		cardiotrophin 1
7894119	0.04474271	-0.51751825	0111	
8156043	0.04480562	-0.33093067	DSAT1	phosphoserine aminotransferase 1
7967660	0.04408187	-0.33108713	RIMBP2	RIMS hinding protein 2
8064502	0.04504240	-0.34076308		small nuclear ribonucleoprotein polypentides R and R1//small nuclear ribonucle
000-002	0.0-00-0-0-0	0.04070500		polypeptides B and B1
8179221	0.04504621	-0.34468993	DPCR1	diffuse panbronchiolitis critical region 1
8122013	0.04529034	-0.43236609	L3MBTL3	I(3)mbt-like 3 (Drosophila)
7896278	0.04530243	-0.31632531		
8103892	0.04534194	-0.31818744		
8039389	0.0454938	-0.53296458	PTPRH	protein tyrosine phosphatase, receptor type, H
7894788	0.04554422	-0.44019559		
8154305	0.04564674	-0.32248636	SELT	selenoprotein T
7964832	0.04565144	-0.51531905		
8139460	0.04567281	-0.30409919	NACAD	NAC alpha domain containing
8080184	0.04572773	-0.43406508	ALAS1	aminolevulinate, delta-, synthase 1
8030749	0.0458141	-0.34302471	-	····,···,··,·,·,
8112327	0.04585459	-0.32325777	CKS1B	CDC28 protein kinase regulatory subunit 1B
8006634	0.04585563	-0.38575586	PIGW	phosphatidylinositol glycan anchor biosynthesis, class W
8144685	0.04588065	-0.46600653		
8028719	0.04588984	-0.30297679	DLL3	delta-like 3 (Drosophila)
8019074	0.04590943	-0.30784858	NPTX1	neuronal pentraxin I
8110227	0.04500510	-0.32463612	LINC5A	unc-5 homolog A (C. elegans)
0110237	0.04033013	0.02403013	SHOUR	

8103399	0.0460248	-0.34168137	PDGFC	platelet derived growth factor C
8040270	0.04604606	-0.30668553	C2orf50	chromosome 2 open reading frame 50
8037502	0.04608252	-0.30936894	NKPD1	NTPase, KAP family P-loop domain containing 1
8138922	0.04609038	-0.43977166	KBTBD2	kelch repeat and BTB (POZ) domain containing 2
7984660	0.04614986	-0.59772748		<u> </u>
8109697	0.04618032	-0.34160251	CCNG1	cyclin G1
8056763	0.04626457	-0.50138702	RPS15	ribosomal protein S15
8175177	0.04644161	-0 40455334	MBNI 3	musclehlind-like splicing regulator 3
7932227	0.04654376	-0 37299962	NMT2	N-myristov/transferase 2
7947649	0.04664147	-0 33839254	CHRM4	cholineraic recentor muscarinic 4
7036016	0.04666062	-0.37132201	EL 140536	El 1/0536 protain
9021470	0.04660602	0.30660604		n E940000 protein
8072002	0.04009090	-0.30000094		
8073992 7003733	0.0407102	-0.31520526		pannexin z
7992732	0.04676655	-0.3732516	20100	
7895246	0.04695187	-0.58685476		
8139828	0.04705725	-0.3096443	14400	
8137931	0.04714298	-0.31127236	MMD2	monocyte to macrophage differentiation-associated 2
8169920	0.04718826	-0.38944057	RBMX2	RNA binding motif protein, X-linked 2
7981783	0.0472862	-0.41832815	UR4N4	oltactory receptor, family 4, subfamily N, member 4
8106962	0.04735951	-0.38628844	ARSK	arylsulfatase family, member K
7940022	0.04739637	-0.43655	RTN4RL2	reticulon 4 receptor-like 2
7961626	0.04744029	-0.66688432	SLCO1A2	solute carrier organic anion transporter family, member 1A2
8175537	0.04757382	-0.49592705	SPANXA2-OT1	SPANXA2 overlapping transcript 1 (non-protein coding)
8074063	0.04763452	-0.39919584	ADM2	adrenomedullin 2
8090639	0.04775727	-0.33510458	PIK3R4	phosphoinositide-3-kinase, regulatory subunit 4
8092095	0.04777602	-0.39039726	TNIK	TRAF2 and NCK interacting kinase
8092314	0.04788819	-0.3010854	DNAJC19	DnaJ (Hsp40) homolog, subfamily C, member 19
7893652	0.04789673	-0.53847207		
7922472	0.04793304	-0.32057647		
7924603	0.04798643	-0.30569892	LBR	lamin B receptor
7945371	0.0480214	-0.31510022	IFITM3	interferon induced transmembrane protein 3
8023710	0.04803044	-0.47750003	CDH19	cadherin 19, type 2
8060418	0.04807919	-0.4299071	SIRPA	signal-regulatory protein alpha
7895267	0.04813092	-0.65876524		
7968670	0.04813242	-0.37148649	UFM1	ubiquitin-fold modifier 1
7988426	0.04815445	-0.37502611	SLC30A4	solute carrier family 30 (zinc transporter), member 4
8162825	0.04823702	-0.43122287		
8056730	0.04828219	-0.34111728	LOC440925	uncharacterized LOC440925
8049435	0.04832363	-0.3644272	SH3BP4	SH3-domain binding protein 4
7925043	0.04833157	-0.36887144	EXOC8///SPRTN	exocyst complex component 8///SprT-like N-terminal domain
8086949	0.04843583	-0.34862771	CCDC51	coiled-coil domain containing 51
8171161	0.04844577	-0.4643664	ARSE	arvlsulfatase E (chondrodysplasia punctata 1)
7995866	0.04845392	-0.30480636	MIR138-2	microRNA 138-2
8080998	0.04845937	-0.42463023		
8110865	0.04848899	-0.31198208	IRX4	iroquois homeobox 4
7992716	0.04856993	-0.4012807	PRSS41	protease serine 41
8100202	0.04863030	-0 59941823	LOC280665///CNGA1	anti-CNG alpha 1 cation channel translation product-like///cyclic nucleotide cate
8033458	0.04865324	-0 3103/500		Ivsonhosholinase II nseudogene////vsonhosholinase II
81/6/27	0.04868057	-0.01004000		notein-L-isoaspartate (D-aspartate) O-methyltransferase domain containing 1
8122026	0.04000337	0.49220402		protein-E-isoaspanale (B-aspanale) O-internymansienase domain containing r
0123330	0.04070114	-0.40407004	INEDD9///INEDD9	expressed, developmentally down-regulated 9
7916120	0.04884725	-0.43140927	TXNDC12	thioredoxin domain containing 12 (endoplasmic reticulum)
7961865	0.04898419	-0.34980162	LYRM5///KRAS	LYR motif containing 5///v-Ki-ras2 Kirsten rat sarcoma viral oncogene homolog
8059731	0.04901827	-0.33575194	PDE6D	phosphodiesterase 6D, cGMP-specific, rod, delta
8003344	0.04902658	-0.33401686	SNAI3	snail family zinc finger 3
7987472	0.04908388	-0.32549584	C15orf56///C15orf56	chromosome 15 open reading frame 56///chromosome 15 open reading frame 5
8131600	0.04909259	-0.82742201	TSPAN13	tetraspanin 13
8052940	0.04913736	-0.34259164	PAIP2B	polv(A) binding protein interacting protein 2B
7896707	0.04927863	-0.396316		
7949843	0.04941305	-0.39256652	CABP2	calcium binding protein 2
8006433	0.04957766	-0.80150668	CCL2	chemokine (C-C motif) ligand 2
7923659	0.04970296	-0.30852482	PPP1R15B	protein phosphatase 1, regulatory subunit 15B
	0.0.010200	5.5500L-TOL		Free Processies in regulatory output in the

7983763	0.04971166	-0.46361643	MAPK6	mitogen-activated protein kinase 6
7941469	0.04977356	-0.36473252	TSGA10IP	testis specific, 10 interacting protein
8169699	0.04985667	-0.33899633		
8016841	0.04990781	-0.41095366	TMEM100	transmembrane protein 100
7902036	0.04997828	-0.34125517		

Comparison of *PTEN* expression in livers between 9 normal and 24 NAFLD/NASH individuals (female).

Probe ID	p-value	Log (Fold Change)	Gene Symbol	Gene Description
7928959	0.88502	-0.06754	PTEN	phosphatase and tensin homolog

miRNA	Gene Name	TargetScan Sites	PicTar Sites
hsa-miR-21-5p	KLHL15	1[18,912]	6[24,6206]
hsa-miR-21-5p	TNPO1	1[6,2277]	6[10,13308]
hsa-miR-21-5p	KLHDC5	1[7,161]	5[9,250]
hsa-miR-21-5p	PLAG1	3[7,512]	5[8,579]
hsa-miR-21-5p	UBE2K	0[0,0]	4[5,104]
hsa-miR-21-5p	YOD1	2[9,3030]	4[11,3172]
hsa-miR-21-5p	HNRNPU	1[5,110]	3[9,114]
hsa-miR-21-5p	CNOT6	1[2,21]	3[8,1002]
hsa-miR-21-5p	RASA1	1[7,212]	3[7,295]
hsa-miR-21-5p	SUZ12	0[0,0]	3[6,1166]
hsa-miR-21-5p	ITGB8	1[4,151]	3[6,1126]
hsa-miR-21-5p	SCML2	1[2,416]	3[5,439]
hsa-miR-21-5p	MAP3K1	1[5,195]	3[5,326]
hsa-miR-21-5p	PURA	0[0,0]	3[5,238]
hsa-miR-21-5p	ACBD5	0[0,0]	3[5,187]
hsa-miR-21-5p	POM121	1[1,1]	3[5,121]
hsa-miR-21-5p	MTAP	1[4,132]	3[4,290]
hsa-miR-21-5p	MTMR12	1[4,35]	3[4,286]
hsa-miR-21-5p	RPS6KA3	1[1,61]	3[4,145]
hsa-miR-21-5p	SLC7A6	1[1,4]	3[3,154]
hsa-miR-21-5p	ZNF367	2[22,8223]	3[22,8379]
hsa-miR-21-5p	RP2	0[0,0]	3[2,53]
hsa-miR-21-5p	TNRC6B	2[15,1113]	3[15,1160]
hsa-miR-21-5p	PURB	1[1,30]	3[12,1575]
hsa-miR-21-5p	CUX1	2[10,536]	3[11,752]
hsa-miR-21-5p	NFIB	1[9,662]	3[11,1028]
hsa-miR-21-5p	LEMD3	1[8,395]	2[9,400]
hsa-miR-21-5p	FRS2	1[1,113]	2[9,392]
hsa-miR-21-5p	YAP1	1[8,191]	2[9,287]
hsa-miR-21-5p	C17orf39	1[1,19]	2[9,227]
hsa-miR-21-5p	SMAD7	1[7,476]	2[8,499]
hsa-miR-21-5p	CPEB3	1[7,562]	2[7,625]
hsa-miR-21-5p	FCHO2	0[0,0]	2[7,533]
hsa-miR-21-5p	PBRM1	1[7,336]	2[7,355]
hsa-miR-21-5p	RMND5A	1[5,1139]	2[7,1393]
hsa-miR-21-5p	DNAJC16	1[3,97]	2[7,106]
hsa-miR-21-5p	AKIRIN1	0[0,0]	2[7,1054]
hsa-miR-21-5p	SPRY2	1[1,35]	2[6,77]
hsa-miR-21-5p	RBPJ	1[1,149]	2[6,686]
hsa-miR-21-5p	ZNF704	0[0,0]	2[6,40]
hsa-miR-21-5p	FAM46A	0[0,0]	2[6,3417]
hsa-miR-21-5p	SATB1	2[6,116]	2[6,116]
hsa-miR-21-5p	PPP1R3B	2[5,64]	2[5,64]

Supplementary Table 2. 219 potential target genes of miR-21

hsa-miR-21-5p	HBP1	0[0,0]	2[5,207]
hsa-miR-21-5p	RNF111	0[0,0]	2[5,189]
hsa-miR-21-5p	PLEKHA1	1[3,514]	2[4,583]
hsa-miR-21-5p	TGFBR2	1[2,3562]	2[4,3696]
hsa-miR-21-5p	EPHA4	1[3,28]	2[4,34]
hsa-miR-21-5p	CSRNP3	1[1,18]	2[4,154]
hsa-miR-21-5p	FBXO11	1[3,1]	2[4,14]
hsa-miR-21-5p	PIK3R1	2[4,108]	2[4,108]
hsa-miR-21-5p	MATR3	0[0,0]	2[3,78]
hsa-miR-21-5p	THRB	2[3,66]	2[3,66]
hsa-miR-21-5p	MKRN1	0[0,0]	2[3,45]
hsa-miR-21-5p	MEF2A	0[0,0]	2[3,279]
hsa-miR-21-5p	POM121C	1[2,7]	2[3,20]
hsa-miR-21-5p	RBMS3	1[2,60]	2[2,68]
hsa-miR-21-5p	IMPAD1	0[0,0]	2[2,66]
hsa-miR-21-5p	STAT3	1[2,168]	2[2,407]
hsa-miR-21-5p	CD47	0[0,0]	2[2,36]
hsa-miR-21-5p	PVRL3	0[0,0]	2[2,354]
hsa-miR-21-5p	ZSWIM6	1[3,395]	2[2,332]
hsa-miR-21-5p	MEF2C	1[1,17]	2[2,25]
hsa-miR-21-5p	RSBN1	0[0,0]	2[2,16]
hsa-miR-21-5p	SOX7	1[2,109]	2[2,109]
hsa-miR-21-5p	GLCCI1	1[16,1845]	2[17,2071]
hsa-miR-21-5p	ZCCHC3	1[10,1917]	2[17,1997]
hsa-miR-21-5p	SKI	1[10,643]	2[13,1587]
hsa-miR-21-5p	ADNP	1[10,455]	2[11,478]
hsa-miR-21-5p	CDC25A	1[10,123]	2[11,292]
hsa-miR-21-5p	GATAD2B	2[10,1633]	2[10,1633]
hsa-miR-21-5p	SYT14	0[0,0]	1[8,333]
hsa-miR-21-5p	BTG2	1[8,2272]	1[8,2272]
hsa-miR-21-5p	FAM63B	1[8,214]	1[8,214]
hsa-miR-21-5p	KIAA1468	0[0,0]	1[7,79]
hsa-miR-21-5p	BTBD3	0[0,0]	1[7,3161]
hsa-miR-21-5p	BCL2	1[7,276]	1[7,276]
hsa-miR-21-5p	E2F3	0[0,0]	1[7,259]
hsa-miR-21-5p	PITX2	1[7,175]	1[7,175]
hsa-miR-21-5p	MSL1	1[7,138]	1[7,138]
hsa-miR-21-5p	ATXN10	0[0,0]	1[6,924]
hsa-miR-21-5p	RAB6A	1[6,64]	1[6,64]
hsa-miR-21-5p	MRPL49	1[6,58]	1[6,58]
hsa-miR-21-5p	SPG20	1[6,286]	1[6,286]
hsa-miR-21-5p	TIMP3	1[6,136]	1[6,136]
hsa-miR-21-5p	KRIT1	1[6,124]	1[6,124]
hsa-miR-21-5p	IL12A	1[6,105]	1[5,96]
hsa-miR-21-5p	UBR3	1[5,93]	1[5,93]
hsa-miR-21-5p	PCBP1	1[5,92]	1[5,92]

hsa-miR-21-5p	TAF5	0[0,0]	1[5,87]
hsa-miR-21-5p	RHOB	1[5,5598]	1[5,5598]
hsa-miR-21-5p	ASF1A	1[5,49]	1[5,49]
hsa-miR-21-5p	SMARCD1	1[5,36]	1[5,36]
hsa-miR-21-5p	CHD7	1[5,3304]	1[5,3304]
hsa-miR-21-5p	LRRC57	1[4,79]	1[4,79]
hsa-miR-21-5p	BCL11A	1[4,43]	1[4,43]
hsa-miR-21-5p	PAN3	1[4,42]	1[4,42]
hsa-miR-21-5p	BRD1	0[0,0]	1[4,163]
hsa-miR-21-5p	TRIM33	1[4,155]	1[4,155]
hsa-miR-21-5p	AP1AR	1[3,75]	1[3,75]
hsa-miR-21-5p	PDZD2	1[3,73]	1[3,73]
hsa-miR-21-5p	SCRN1	1[3,70]	1[3,70]
hsa-miR-21-5p	NFIA	1[3,55]	1[3,55]
hsa-miR-21-5p	GLYR1	1[3,428]	1[3,428]
hsa-miR-21-5p	ALX1	1[3,33]	1[3,33]
hsa-miR-21-5p	ZDHHC17	0[0,0]	1[3,178]
hsa-miR-21-5p	RECK	1[3,161]	1[3,161]
hsa-miR-21-5p	DLX2	1[3,146]	1[3,146]
hsa-miR-21-5p	KIAA2026	0[0,0]	1[3,108]
hsa-miR-21-5p	CBX4	1[22,2569]	1[22,2569]
hsa-miR-21-5p	SPIN1	0[0,0]	1[2,94]
hsa-miR-21-5p	TGFBI	1[2,930]	1[2,930]
hsa-miR-21-5p	MBTPS2	0[0,0]	1[2,8]
hsa-miR-21-5p	PPP3CA	1[2,497]	1[2,497]
hsa-miR-21-5p	PER2	1[2,484]	1[2,484]
hsa-miR-21-5p	C11orf95	1[2,48]	1[2,48]
hsa-miR-21-5p	PPARA	1[2,40]	1[2,40]
hsa-miR-21-5p	JHDM1D	1[2,39]	1[2,39]
hsa-miR-21-5p	SOS2	0[0,0]	1[2,33]
hsa-miR-21-5p	SLC9A6	0[0,0]	1[2,288]
hsa-miR-21-5p	PCSK6	1[2,232]	1[2,232]
hsa-miR-21-5p	EHD1	1[2,228]	1[2,228]
hsa-miR-21-5p	TESK2	0[0,0]	1[2,20]
hsa-miR-21-5p	PCBP2	1[2,171]	1[2,171]
hsa-miR-21-5p	PSRC1	0[0,0]	1[2,171]
hsa-miR-21-5p	CRIM1	0[0,0]	1[2,1621]
hsa-miR-21-5p	KLF6	1[2,137]	1[2,137]
hsa-miR-21-5p	WWP1	1[2,132]	1[2,132]
hsa-miR-21-5p	ARHGAP24	1[2,13]	1[2,13]
hsa-miR-21-5p	TSHZ1	0[0,0]	1[2,115]
hsa-miR-21-5p	UBE2D3	1[2,105]	1[2,105]
hsa-miR-21-5p	STK40	1[2,0]	1[2,0]
hsa-miR-21-5p	XKR6	1[2,0]	1[2,0]
hsa-miR-21-5p	PELI1	1[19,4917]	1[19,4917]
hsa-miR-21-5p	EIF2C2	0[0,0]	1[19,1480]

hsa-miR-21-5p	PDCD4	1[18,1672]	1[18,1672]
hsa-miR-21-5p	MBNL1	1[15,3279]	1[15,3279]
hsa-miR-21-5p	STAG2	1[14,394]	1[14,394]
hsa-miR-21-5p	KLF3	1[11,4344]	1[11,4344]
hsa-miR-21-5p	EIF1AX	1[10,269]	1[10,269]
hsa-miR-21-5p	DAZL	0[0,0]	1[1,9]
hsa-miR-21-5p	NAA50	0[0,0]	1[1,6]
hsa-miR-21-5p	RASGRP1	1[1,5]	1[1,5]
hsa-miR-21-5p	SETD1B	1[1,47]	1[1,47]
hsa-miR-21-5p	NIPBL	0[0,0]	1[1,212]
hsa-miR-21-5p	SSFA2	0[0,0]	1[1,1996]
hsa-miR-21-5p	ELF2	1[1,19]	1[1,19]
hsa-miR-21-5p	MATN2	1[2,30]	1[1,17]
hsa-miR-21-5p	GLIS2	1[1,125]	1[1,125]
hsa-miR-21-5p	JPH1	1[1,12]	1[1,12]
hsa-miR-21-5p	MAPRE1	0[0,0]	1[1,11]
hsa-miR-21-5p	VCL	2[7,576]	0[0,0]
hsa-miR-21-5p	RAB22A	2[6,2476]	0[0,0]
hsa-miR-21-5p	PTPN9	1[9,425]	0[0,0]
hsa-miR-21-5p	FBXO28	1[9,137]	0[0,0]
hsa-miR-21-5p	ZNF217	1[8,1458]	0[0,0]
hsa-miR-21-5p	BRWD1	1[7,682]	0[0,0]
hsa-miR-21-5p	GNG12	1[6,587]	0[0,0]
hsa-miR-21-5p	LRP6	1[6,345]	0[0,0]
hsa-miR-21-5p	PTPN14	1[6,1660]	0[0,0]
hsa-miR-21-5p	ZFP36L2	1[6,115]	0[0,0]
hsa-miR-21-5p	DNAJA2	1[5,96]	0[0,0]
hsa-miR-21-5p	GAB1	1[5,831]	0[0,0]
hsa-miR-21-5p	CHIC1	1[5,50]	0[0,0]
hsa-miR-21-5p	SC5DL	1[5,245]	0[0,0]
hsa-miR-21-5p	SESTD1	1[5,164]	0[0,0]
hsa-miR-21-5p	DCUN1D3	1[4,69]	0[0,0]
hsa-miR-21-5p	GTPBP1	1[4,62]	0[0,0]
hsa-miR-21-5p	PRKCE	1[4,45]	0[0,0]
hsa-miR-21-5p	MAP2K3	1[4,37]	0[0,0]
hsa-miR-21-5p	PFKM	1[4,336]	0[0,0]
hsa-miR-21-5p	EIF4EBP2	1[4,303]	0[0,0]
hsa-miR-21-5p	SNTB2	1[4,214]	0[0,0]
hsa-miR-21-5p	RFFL	1[3,84]	0[0,0]
hsa-miR-21-5p	EIF2C4	1[3,6]	0[0,0]
hsa-miR-21-5p	PAG1	1[3,50]	0[0,0]
hsa-miR-21-5p	PGRMC2	1[3,469]	0[0,0]
hsa-miR-21-5p	S100A10	1[3,381]	0[0,0]
hsa-miR-21-5p	GXYLT1	1[3,336]	0[0,0]
hsa-miR-21-5p	LIFR	1[3,3256]	0[0,0]
hsa-miR-21-5p	FOXP2	1[3,32]	0[0,0]

hsa-miR-21-5p	5-Mar	1[3,14]	0[0,0]
hsa-miR-21-5p	BCL7A	1[21,973]	0[0,0]
hsa-miR-21-5p	UBN2	1[2,99]	0[0,0]
hsa-miR-21-5p	TNFRSF11B	1[2,91]	0[0,0]
hsa-miR-21-5p	KLF5	1[2,823]	0[0,0]
hsa-miR-21-5p	ARMCX1	1[2,78]	0[0,0]
hsa-miR-21-5p	DUSP8	1[2,66]	0[0,0]
hsa-miR-21-5p	TOPORS	1[2,61]	0[0,0]
hsa-miR-21-5p	TET1	1[2,38]	0[0,0]
hsa-miR-21-5p	AIM1L	1[2,37]	0[0,0]
hsa-miR-21-5p	MBLAC2	1[2,35]	0[0,0]
hsa-miR-21-5p	MSH2	1[2,325]	0[0,0]
hsa-miR-21-5p	SLC10A7	1[2,26]	0[0,0]
hsa-miR-21-5p	COL4A1	1[2,2419]	0[0,0]
hsa-miR-21-5p	GPR64	1[2,24]	0[0,0]
hsa-miR-21-5p	BMPR2	1[2,210]	0[0,0]
hsa-miR-21-5p	ZBTB47	1[2,2042]	0[0,0]
hsa-miR-21-5p	MCTP2	1[2,20]	0[0,0]
hsa-miR-21-5p	C5orf41	1[2,137]	0[0,0]
hsa-miR-21-5p	SESN1	1[2,132]	0[0,0]
hsa-miR-21-5p	C17orf75	1[2,13]	0[0,0]
hsa-miR-21-5p	FAM3C	1[2,12]	0[0,0]
hsa-miR-21-5p	ARHGEF12	1[2,11]	0[0,0]
hsa-miR-21-5p	CCR7	1[2,108]	0[0,0]
hsa-miR-21-5p	C10orf12	1[2,10]	0[0,0]
hsa-miR-21-5p	BRD2	1[17,667]	0[0,0]
hsa-miR-21-5p	CDK6	1[13,1469]	0[0,0]
hsa-miR-21-5p	KBTBD6	1[11,635]	0[0,0]
hsa-miR-21-5p	ARHGAP31	1[1,8]	0[0,0]
hsa-miR-21-5p	FAM126B	1[1,70]	0[0,0]
hsa-miR-21-5p	HSDL2	1[1,6]	0[0,0]
hsa-miR-21-5p	MBNL3	1[1,5]	0[0,0]
hsa-miR-21-5p	DAG1	1[1,213]	0[0,0]
hsa-miR-21-5p	PIKFYVE	1[1,21]	0[0,0]
hsa-miR-21-5p	TNS1	1[1,189]	0[0,0]
hsa-miR-21-5p	RALGPS2	1[1,13]	0[0,0]
hsa-miR-21-5p	FAM13A	1[1,13]	0[0,0]
hsa-miR-21-5p	LMBR1	1[1,107]	0[0,0]

P Value	Log FC/Fatty liver/Normal)	Gene symbol	Gene Name	miRNA	TargetScan Sites	PicTar Sites
0.01630449	-0.54013801	KLHL15	kelch-like family member 15	hsa-miR-21-5p	1[18,912]	6[24,6206]
0.04430029	-0.37042566		cytoplasmic polyadenylation element binding protein 3	hsa-miR-21-5p	1[7,562]	2[7,625]
0.03034044	-0.31292391	HBF I		nsa-mik-21-5p	0[0,0]	2[0,207]
0.02570623	-0.39409707	IMPAD1	inositol monophosphatase domain containing 1	hsa-miR-21-5p	0[0,0]	2[2,66]
0.00491062	-0.34885596	PVRL3	poliovirus receptor-related 3	nsa-miR-21-5p	0[0,0]	2[2,354]
0.00865386	-0.43339143	SOX7	SRY (sex determining region Y)-box 7	hsa-miR-21-5p	1[2,109]	2[2,109]
0.00260512	-0.71029488	RHOB	ras homolog family member B	hsa-miR-21-5p	1[5,5598]	1[5,5598]
0.03194318	-0.3484977	AP1AR	adaptor-related protein complex 1 associated regulatory protein	hsa-miR-21-5p	1[3,75]	1[3,75]
0.00897159	-0.31889815	MBTPS2	membrane-bound transcription factor peptidase, site 2	hsa-miR-21-5p	0[0,0]	1[2,8]
0.01321711	-1.08633018	KLF6	Kruppel-like factor 6	hsa-miR-21-5p	1[2,137]	1[2,137]
0.02377635	-0.44395978	TNFRSF11B	tumor necrosis factor receptor superfamily, member 11b	hsa-miR-21-5p	1[2,91]	0[0,0]
0.03118009	-0.75072228	KLF5	Kruppel-like factor 5 (intestinal)	hsa-miR-21-5p	1[2,823]	0[0,0]
0.04644161	-0.40455334	MBNL3	muscleblind-like splicing regulator 3	hsa-miR-21-5p	1[1,5]	0[0,0]

Supplementary Table 3. Thirteen down-regulated genes in livers of human NAFLD/NASH that have binding motifs for miR-21.

Supplementary Table 4. qRT-PCR primers used in this study

Gene Name	Forward primer	Reverse primer
Human HBP1	GAGTGAACCAGCCTTCCCTCAT	ACAGTGCCAGACAGTTGAAGGC
Moue Hbp1	TGGCTTGCTCACTGTAGAGTGC	CAGGAGGTAGACATACGTCACC
Human <i>p5</i> 3	CCTCAGCATCTTATCCGAGTGG	TGGATGGTGGTACAGTCAGAGC
Mouse p53	CACGTACTCTCCTCCCCTCAAT	AACTGCACAGGGCACGTCTT
Human SREBP1C	ACTTCTGGAGGCATCGCAAGCA	AGGTTCCAGAGGAGGCTACAAG
Mouse Srebp1c	CGACTACATCCGCTTCTTGCAG	CCTCCATAGACACATCTGTGCC
Human FASN	TTCTACGGCTCCACGCTCTTCC	GAAGAGTCTTCGTCAGCCAGGA
Mouse Fasn	CACAGTGCTCAAAGGACATGCC	CACCAGGTGTAGTGCCTTCCTC
Human SCD1	CCTGGTTTCACTTGGAGCTGTG	TGTGGTGAAGTTGATGTGCCAGC
Mouse Scd1	GCAAGCTCTACACCTGCCTCTT	CGTGCCTTGTAAGTTCTGTGGC
Human GPAT	TTGTGGCTTGCCTGCTCCTCTA	AATCACGAGCCAGGACTTCCTC
Mouse Gpat	GCAAGCACTGTTACCAGCGATC	TGCAATCAGCCTTCGTCGGAAG
Human CCND1	TCTACACCGACAACTCCATCCG	TCTGGCATTTTGGAGAGGAAGTG
Mouse Ccnd1	GCAGAAGGAGATTGTGCCATCC	AGGAAGCGGTCCAGGTAGTTCA
Human CCNB1	GACCTGTGTCAGGCTTTCTCTG	GGTATTTTGGTCTGACTGCTTGC
Mouse Ccnb1	AGAGGTGGAACTTGCTGAGCCT	GCACATCCAGATGTTTCCATCGG

	0	Ċt	۵۵	Ct	2-2	,Ct	(Ċt	Δ	Ct	2-	∆Ct	Ave.	2 ⁻ Δ ^{Ct}	Fold Change
miRNA ID	WT1	WT2	WT1	WT2	WT1	WT2	DKO1	DKO2	DKO1	DKO2	DKO1	DKO2	WT	DKO	DKO/WT
mmu-miR-21	25.504	23.928	8.516	6.348	0.00273167	0.0122761	33.838	26.752	17.688	11.300	4.7357E-06	0.00039661	0.007503905	0.000200672	0.026742
mmu-miR-122	22.365	21.345	5.377	3.765	0.02406367	0.0735567	26.180	24.536	10.030	9.084	0.00095646	0.00184265	0.048810186	0.001399559	0.028674
mmu-miR-192	23.378	22.054	6.390	4.474	0.0119239	0.0449978	26.303	24.690	10.153	9.238	0.0008783	0.00165609	0.028460862	0.001267196	0.044524
mmu-miR-30c	23.793	22.661	6.805	5.081	0.00894316	0.0295438	26.099	25.122	9.949	9.670	0.0010117	0.00122755	0.019243496	0.001119627	0.058182
hsa-miR-30a- 3p	23.769	24.231	6.781	6.651	0.00909318	0.0099506	25.672	26.853	9.522	11.401	0.00136017	0.00036979	0.00952189	0.000864979	0.090841
mmu-miR-19b	24.509	22.691	7.521	5.111	0.00544444	0.0289358	27.076	24.557	10.926	9.105	0.00051398	0.00181603	0.017190126	0.001165003	0.067772
mmu-miR-30b	24.212	23.308	7.224	5.728	0.00668897	0.0188669	26.802	25.150	10.652	9.698	0.00062148	0.00120396	0.012777924	0.00091272	0.071429
mmu-let-7g	26.364	25.287	9.376	7.707	0.00150502	0.0047859	28.757	27.006	12.607	11.554	0.00016029	0.00033258	0.00314545	0.000246438	0.078347
mmu-miR-30e	25.973	24.847	8.985	7.267	0.00197354	0.0064925	27.878	27.017	11.728	11.565	0.0002948	0.00033006	0.004233042	0.000312426	0.073806
mmu-miR-29a	24.430	22.458	7.442	4.878	0.00575088	0.0340076	26.324	24.590	10.174	9.138	0.00086561	0.00177496	0.019879228	0.001320282	0.066415
mmu-miR-26b	26.134	24.538	9.146	6.958	0.00176514	0.0080433	28.187	26.264	12.037	10.812	0.00023796	0.00055624	0.004904211	0.0003971	0.080971
mmu-miR-26a	25.490	23.226	8.502	5.646	0.00275831	0.0199703	27.288	24.873	11.138	9.421	0.00044374	0.0014588	0.011364307	0.000951271	0.083707
mmu-miR-17	25.260	24.397	8.272	6.817	0.00323504	0.0088691	26.815	25.722	10.665	10.270	0.00061591	0.00080988	0.006052062	0.000712895	0.117794
mmu-miR-106a	26.483	25.696	9.495	8.116	0.00138586	0.0036045	27.867	27.027	11.717	11.575	0.00029705	0.00032778	0.002495164	0.000312414	0.125208
mmu-miR-484	25.192	25.251	8.204	7.671	0.00339117	0.0049068	26.622	26.389	10.472	10.937	0.00070407	0.00051008	0.004148991	0.000607072	0.146318
mmu-miR- 125b-5p	27.341	26.233	10.353	8.653	0.0007646	0.0024842	28.758	27.099	12.608	11.647	0.00016018	0.00031182	0.001624405	0.000236001	0.145285
mmu-miR-191	23.274	23.119	6.286	5.539	0.0128152	0.0215078	24.210	24.156	8.060	8.704	0.00374713	0.00239792	0.017161476	0.003072524	0.179036
mmu-miR-24	23.695	22.427	6.707	4.847	0.00957176	0.0347462	24.886	22.981	8.736	7.529	0.00234532	0.00541433	0.022158993	0.003879826	0.17509
mmu-miR-16	22.631	22.048	5.643	4.468	0.02001187	0.0451854	23.136	22.835	6.986	7.383	0.00788868	0.00599095	0.03259862	0.006939818	0.212887
mmu-miR-2134	16.977	18.260	-0.011	0.680	1.00765426	0.6241652	17.246	18.971	1.096	3.519	0.46781165	0.08723188	0.815909718	0.277521769	0.340138
mmu-miR-2146	20.563	21.864	3.575	4.284	0.0839108	0.0513319	21.240	22.102	5.090	6.650	0.02936009	0.00995751	0.067621352	0.0196588	0.290719
mmu-miR-142- 3p	28.469	27.431	11.481	9.851	0.00034984	0.0010828	29.132	27.100	12.982	11.648	0.0001236	0.0003116	0.000716329	0.000217603	0.303776
mmu-miR-126- 3p	21.265	20.774	4.277	3.194	0.05158161	0.1092723	21.934	20.262	5.784	4.810	0.01814857	0.03564889	0.080426962	0.026898731	0.334449
mmu-miR-139- 5p	27.021	26.326	10.033	8.746	0.00095448	0.0023291	27.314	26.078	11.164	10.626	0.00043581	0.00063278	0.001641798	0.000534299	0.325435

Supplementary Table 5. miRNAs that are highly and specifically expressed in hepatocytes.

mmu-miR- 1937b	18.374	18.697	1.386	1.117	0.38262405	0.4610514	18.818	18.287	2.668	2.835	0.15734456	0.1401457	0.421837722	0.148745128	0.352612
mmu-miR-145	27.880	25.955	10.892	8.375	0.00052624	0.0030121	28.442	25.423	12.292	9.971	0.00019941	0.00099639	0.001769184	0.000597899	0.337952
mmu-miR-486	26.424	26.279	9.436	8.699	0.00144371	0.0024062	26.161	26.332	10.011	10.880	0.00096915	0.00053063	0.00192498	0.000749889	0.389557
mmu-miR-126- 5p	25.840	24.903	8.852	7.323	0.00216413	0.0062454	26.368	24.118	10.218	8.666	0.00083961	0.00246192	0.004204742	0.001650763	0.392596
mmu-miR-805	23.583	24.462	6.595	6.882	0.01034444	0.0084784	23.665	23.949	7.515	8.497	0.00546713	0.00276789	0.009411399	0.004117508	0.437502
mmu-miR-146a	25.255	24.807	8.267	7.227	0.00324627	0.0066751	25.608	24.004	9.458	8.552	0.00142186	0.00266435	0.004960677	0.002043108	0.411861
mmu-miR- 1937c	21.345	21.400	4.357	3.820	0.04879918	0.0708053	21.443	20.826	5.293	5.374	0.02550633	0.02411375	0.05980222	0.02481004	0.414868
mmu-miR-721	22.005	24.120	5.017	6.540	0.03088394	0.0107464	23.009	22.368	6.859	6.916	0.0086146	0.00828088	0.020815179	0.00844774	0.405845
mmu-miR-223	27.620	26.374	10.632	8.794	0.00063016	0.0022529	27.790	25.456	11.640	10.004	0.00031334	0.00097386	0.001441529	0.000643598	0.446469
mmu-miR-150	27.367	27.284	10.379	9.704	0.00075095	0.001199	27.315	26.413	11.165	10.961	0.00043551	0.00050166	0.000974954	0.000468586	0.480624

Supplementary Table 6. Comparison of mice treated with miR-21-ASO or miR-21-MM-ASO for 4 weeks

Parameter	miR-21-MM-ASO treated mice	miR-21-ASO treated mice
Body weight (g)	34±3	32±4
Liver weight	1.9±0.3	1.7±0.5
Liver cholesterol concentration (mg/g)	3.8±0.7	4.0±0.7
Total plasma cholesterol(mg/dl)	140±19	129±16.2
Total plasma triglycerides (mg/dl)	113±20.3	105±10.3

Supplementary Table 7. Comparison of mice treated with miR-21-ASO or miR-21-MM-ASO for 8 weeks

Parameter	miR-21-MM-ASO treated mice	miR-21-ASO treated mice
Body weight (g)	43±4	41±3.4
Liver weight (g)	2.4±0.4	1.9±0.38*
Liver cholesterol concentration (mg/g)	4.5±0.6	4.3±0.5
Liver triglyceride concentration (mg/g)	142±16	75±8.1*
Total plasma cholesterol(mg/dl)	191±21	179±14.6
Total plasma triglycerides (mg/dl)	143±15.2	136±9.8

	ASO treatment					
ASO	(weeks)	Albumin (g/dL)	Total Bilirubin (mg/dL)	ALP (U/L)	ALT (U/L)	AST (U/L)
miR-21-MM-ASO (control)	4	3.21+0.15	0.37+0.08	56.8+9.3	52.1+2.4	326.6+19.7
miR-21-ASO	4	3.28+0.18	0.31+0.14	45.0+15.8	40.7+2.3*	261.8+23.8*
miR-21-MM-ASO (control)	8	3.45+0.17	0.28+0.03	74.75+6.7	55.3+7.8	337.8+30.8
miR-21-ASO	8	3.61+0.29	0.23+0.05	58.6+23.27	48.8+5.60	231.16+24.1*

Supplementary Table 8. Comparison of serum chemistries of mice treated with miR-21-ASO or miR-21-MM-ASO

Table 1. 1219 down-regulated probes in livers of human patients with NAFLD/NASH.

Table 2. 219 potential target genes of miR-21.

Table 3. Thirteen down-regulated genes in livers of human NAFLD/NASH that have binding motifs for miR-21.

Table 4. RT-PCR primers used in this study

Table 5. miRNAs that are highly expressed in hepatocytes. Highly-expressed miRNAs were identified by comparing miRNA profiles of livers between wild-type and *Dicer1* knockout mice. Internal control is U6 nuclear small RNA and U6 was measured five times in each sample. The average Ct value of U6: WT1: 16.998; WT2: 17.58; DKO1: 16.150; and DKO2: 15.452. WT: Wild type; and DKO: *Dicer1* knockout.

Table 6. Comparison of mice treated with miR-21-ASO and miR-21-MM-ASO for four weeks.Data represent mean \pm SD. Mann-Whitney was used for statistical analysis.

Table 7. Comparison of mice treated with miR-21-ASO and miR-21-MM-ASO for eight weeks. Data represent mean \pm SD. Mann-Whitney was used for statistical analysis. *p < 0.05.

Table 8. Comparison of serum chemistry of mice treated with miR-21-ASO and miR-21-MM-ASO. Data represent mean \pm SD. Mann-Whitney was used for statistical analysis. *p < 0.05.