ROLE OF MICRORNA LET-7 IN PANCREATIC BETA CELLS

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ROLE OF MICRORNA LET-7 IN PANCREATIC BETA CELLS

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List of abbreviations

AGO2: Argonaute

EIF4EBP2: Eukaryotic translation initiation factor 4E binding protein 2

IGF1R: Insulin-like growth factor 1 receptor

INSR/IR: Insulin receptor

IRS-1/2: Insulin receptor substrate

ITT/GTT: Insulin tolerance tests/Glucose tolerance tests

GLUT: Glucose transporter

GSIS: Glucose stimulated insulin secretion

HMGA2: High mobility group AT-hook 2

MAP kinase: Mitogen-activated protein kinase

MODY genes: Maturity onset diabetes of youth genes

mTOR: mammalian target of rapamycin

MTPN: mytrophin

PDX-1: Pancreatic and duodenal homeobox 1

PI3 kinase: Phosphoinositide 3 kinase

PKB/Akt: protein kinase B

RISC: RNA-induced silencing complex

STTM: Short tandem target mimic

T1D: Type 1 diabetes

T2D: Type 2 diabetes

TNF-a: Tumor necrosis factor alpha

Abstract
MicroRNAs (miRNAs) are small non-coding RNAs that inhibit gene expression at transcriptional or post-transcriptional level. Let-7 family is among the first identified human miRNAs and regulates multiple cellular processes including glucose metabolism in multiple organs. It has been reported that overexpression of let-7 resulted in insulin resistance and impaired glucose tolerance through repressing insulin signaling pathway in both muscle and liver. However, the role and mechanism underlying let-7 function in pancreatic beta-cells have yet to be elucidated.

Let-7 family contains nine members, which poses a significant challenge in complete deletion of this miRNA family. To study the function of let-7 and to overcome the functional redundancies of various let-7 members in pancreatic beta-cells, the highly expressed let-7a and let-7b were blocked simultaneously using short tandem target mimic (STTM) approach developed in our laboratory. Introducing STTM-let7 into beta-cells markedly increased the expression of Caspase 3, a direct target of let-7, confirming a sufficient functional knockdown of let-7a/b by STTM-let7. STTM-let7 enhanced apoptotic cell death induced by cytokine, indicating that let-7a/b is able to protect from apoptosis through attenuating Caspase 3 expression in pancreatic beta-cells. In contrast to the previous observation that let-7 silencing increases insulin signaling in muscle and liver, inhibition of let-7 with STTM-let7 significantly repressed glucose-stimulated insulin signaling in pancreatic beta-cells, leading to impaired insulin secretion and reduced beta-cell proliferation. Taken together, an appropriate level of let-7 is essential in maintaining beta-cell function and viability. Dysregulation of let-7 may contribute to the pathogenesis of type2 diabetes.
1. Introduction

1.1 Diabetes and pancreatic beta-cells

Diabetes is a disorder of metabolism in which a person has high blood glucose level (hyperglycemia). If left untreated, hyperglycemia can cause serious complications, such as heart diseases and kidney failure(1). Diabetes is due to either insufficient amount of insulin or dysfunction of insulin effects on the body. There are three types of diabetes: type 1 diabetes, type 2 diabetes and gestational diabetes. Type 1 diabetes (T1D) is characterized by loss of insulin-producing pancreatic 1-cell caused by autoimmune attack on the 1-cells. Type 1 diabetes is usually diagnosed in children and young adults and previously called juvenile diabetes or "insulin-dependent diabetes"(2). Type 2 Diabetes (T2D) results from the combination of resistance to insulin action in muscle and adipocytes and insufficient insulin production 1-cells. Approximately 90% of diabetic patients are type 2 diabetes(3). Gestational diabetes is a type of diabetes that only occurs during pregnancy in females who previously did not have diabetes.

Type 2 Diabetes is one of the most prevalent diseases around the world. There were approximately 285 million people diagnosed with T2D in 2010(4). Development of type 2 diabetes involves multiple metabolic defects, mainly due to insulin resistance and 1-cell dysfunction. Most people with insulin resistance fail to respond to normal circulating insulin, causing reduced glucose uptake in muscle and fat tissues, increased glucose production and release in liver, which all in turn result in elevated blood glucose levels(5). If insulin resistance exists, much higher insulin requires to be secreted to blood from pancreatic 1-cells, a condition called hyperinsulinemia(6). Hyperglycemia combined with hyperinsulinemia generate various toxicities, such as free fatty acids, reactive oxygen species and inflammatory cytokines, which result in 1-cell dysfunction including reduced insulin secretion and insulin biosynthesis, increased 1-cell death and eventually loss of 1-cell mass(7-9).

Pancreatic 1-cells are the only source for producing insulin and insulin is the key hormone responsible for maintaining glucose homeostasis. In 1-cell, insulin
biosynthesis and release are tightly regulated by insulin signaling pathway that maintain the 1-cell fate and activate specific transcription factors in response to the change of plasma glucose(10). The insulin signaling pathway involves insulin activated insulin receptor (IR), insulin receptor substrate (IRS1/2), phosphorylation, and activation of phosphatidylinositol 3-kinase (PI3-K), Akt (also known as protein kinase B), and mammalian target of rapamycin (mTOR)(Figure 1.1)(10). Studies in diabetic animal models and humans have consistently demonstrated that dysregulation of insulin signaling is associated with impaired insulin secretion and insulin biosynthesis. Overexpression of insulin receptor in 1-cells was found to promote insulin transcription and regulate the steady-state insulin content (11). Leibiger and his colleagues reported type A insulin receptor controlled insulin gene transcription whereas stimulation of type B insulin receptor promoted 1-cell glucokinase gene expression (12). Insulin receptor substrate 2 (IRS-2) branch of the insulin/insulin-like growth factor signaling is capable of mediating pancreatic 1-cell proliferation and function. Exendin-4 promoted IRS-2 expression and Akt phosphorylation were able to delay progression of diabetes and to stimulate insulin secretion (13). IRS-2 was also shown to be regulated by negative feedback effects of mammalian target of rapamycin (mTOR). Chronic activation of mTOR resulted in degradation of IRS-2 followed by deactivation of Akt/PKB and elevation of 1-cell apoptosis.(14) This work provided another mechanism indicating why 1-cell is damaged by chronic hyperglycemia in the development of type 2 diabetes.
Figure 1.1 Glucose and insulin stimulated insulin signaling pathways in pancreatic 1-cells. Insulin molecules interact with Insulin receptor (IR) and activate both the mitogenic (via MAP kinase) and metabolic branches involving phosphoinositide 3 (PI3)-kinase, protein kinase B (PKB, also called Akt), mammalian target of rapamycin (mTOR). Adapted from (10).

The development of type 2 diabetes is associated with a loss of 1-cell mass. The 1-cell mass is maintained by the dynamic balance of proliferation and cell death. Under normal conditions, the proliferation and apoptosis rate are very low(15). However, in the late phase of diabetes, 1-cell mass is significantly decreasing due to an enhanced 1-cell apoptosis. Evidences have demonstrated the proinflammatory cytokines stimulate 1-cell apoptosis by activating Bcl-2 regulated intrinsic apoptotic pathway(16)(Figure 1.2). The proapoptotic members, Bax and Bak, directly promote mitochondrial swelling and release of cytochrome C. Cytochrome C release leads to the formation of an apoptosome, which in turn activates caspase-3 and ultimately induces cell apoptosis(17,18). Antiapoptotic Bcl-2 family sequesters Bax and Bak in the cytosol, thus inhibiting apoptosis. Understanding how cytokine triggers 1-cell apoptosis is likely to shed new light
on mechanisms of 1-cell loss in diabetes and may therefore find the way to improve therapeutic intervention.

**Figure 1.2 Diagram of cell death signaling.** Cell death can be initiated through intrinsic pathway (Mitochondria) or extrinsic pathway (ligand-receptor binding). Both of them require the activation of initiator caspases (CASP8, CASP9) and effector caspases (CASP3). Let-7 was revealed to suppress the expression of Caspase-3 (19). Bax and Bcl-2 are antagonistic regulators mediate the release of Cytochrome C from mitochondria.

**1.2 Role of microRNA in beta cells**

MicroRNA (miRNA) is a small (-22 nucleotides) non-coding RNA located in the introns or non-coding region of genome. MiRNAs can be found in plants, animals and some virus, in which they regulate gene expression on transcriptional or post-transcriptional level. Containing the complementary sequences to the messenger RNA (mRNA), miRNAs interact with their target mRNA through base pairing, resulting in inhibition or degradation of the mRNA(20). In human body, miRNAs mainly repress translation by binding imperfectly with 3’-untranslated region of their targets. Generally, one single miRNA regulates expression of multiple genes and one single mRNA is controlled by a combination of multiple
miRNAs(20,21). In animal cells, the biogenesis of miRNAs starts from transcribing miRNA genes to primary transcripts (pri-miRNAs) by RNA polymerase II (Pol II). The poly-A tail of pri-miRNA is cropped by the Drosha complex, generating the precursor miRNA (pre-miRNA) which has a hairpin structure with a -2-nucleotide 3' overhang. Pre-miRNA is recognized by exportin-5 and transported into cytoplasm in which the pre-miRNA is cleaved by Dicer to produce miRNA duplex. Dicer and Argonaute (AGO2)(22) mediate the processing of pre-miRNA and assemble the RNA-induced silencing complex (RISC). One strand of the miRNA is degraded whereas the other is incorporated into AGO protein as the mature miRNA. AGO2 is responsible for possessing the inhibition of target mRNA(23,24) (Figure 1.3).

**Figure 1.3 Diagram of miRNA biogenesis in animals.** miRNAs are first transcribed into pri-miRNA and undergone the processing of Drosha. After exportation out of nucleus, Dicer, TAR RNA-binding protein (TARBP2), protein activator of the interferon induced protein kinase (PACT) and Argonaute (AGO2)
mediate the cleavage of pre-miRNA and are assembled to RNA-induced silencing complex (RISC). Adapted from (24).

MicroRNAs play diverse roles in many biological processes, including proliferation, differentiation and apoptosis. Dysregulations of miRNAs thus are involved in various diseases like cancer and diabetes. Several miRNAs have been identified as pathological factors contribute to the development of diabetes (25). Previously, miR-375 was found to be elevated in patients with type 2 diabetes. Overexpression of miR-375 results in reduced expression of mytrophin (MTPN) and t-SNAREs yeast homologue 1A (Vt1a), leading to impairment of insulin secretion (26,27). MiR-375 was further found to be required for maintain 1-cell mass. Genetic deletion of miR-375 impairs the proliferation of pancreas and results in a severely diabetic state, whereas the miR-375 knockout mice have increased pancreatic alpha cell mass and glucagon levels (28). miR-30d was shown to associate with the expression of MafA, an insulin gene transcription factors as well as prevent the reduction of IRS-2 from tumor necrosis factor alpha (TNF-a) exposure (29). MiR-29a/b/c was increased in isolated islets of prediabetic non-obese diabetic (30) mice with impaired insulin secretion (31). Other miRNAs regulating pancreatic 1-cell function include miR-144 that was upregulated in blood, pancreas, liver and skeletal muscles of type 2 diabetes model rats (32,33) and miR-24 that overexpression inhibited insulin secretion and 1-cell proliferation by targeting two maturity onset diabetes of youth (MODY) genes (34).

1.3 Let-7 family of microRNAs

In this study, we identified the specific functions of a highly conserved miRNA family, let-7, in pancreatic 1-cells. Let-7 family of miRNAs is one of the most abundant miRNAs expressed among animals. In mice, there are nine different let-7 members encoded by 12 genes in which let-7a has identical sequence across various species from C. elegans to human. All of let-7 family members are believed to have similar functions because of the same seed sequences in their mature form (35,36). Let-7 family was first identified as tumor suppressor by negatively regulating many oncogenes. In human cancers, loss of let-7 was
discovered resulting in upregulation of some cell-cycle factors, including RAS, MYC, HMGA2 (high mobility group AT-hook 2), cyclin D and CDC34. Increasing of these let-7-targeted genes leads to dysfunction of cell growth and proliferation and termination of normal differentiation and the emergence of malignancy(37-39). Interestingly, let-7a was nevertheless found to suppress cytokine-induced cancer cell death by inhibiting the expression of caspase-3, an executioner caspase that initiate apoptosis (19) (Figure 1.2). Let-7 family also regulated glucose metabolism in multiple organ through lin28/let-7 axis. Lin28 and let-7 were mutually antagonistic regulators involved in glucose homeostasis. Lin28 transgenic mice were resistant to obesity and diabetes and exhibited enhanced glucose tolerance. Hao Zhu and his colleagues reported that proteins associated with insulin signaling, such as IRS-2, IGF1R (insulin-like growth factor 1 receptor), PIK3IP1 (phosphoinositide-3-kinase interacting protein 1), Akt2 and EIF4EBP2 (eukaryotic translation initiation factor 4E-binding protein 2) were all predicted to contain let-7 binding sites(30). Further study indicated transgenic mice globally overexpress let-7 exhibited glucose intolerance and reduced glucose-stimulated insulin secretion. Glucose tolerance tests (GTT) and insulin tolerance tests (ITT) in transgenic mice suggested insulin resistance in peripheral tissues and the reduction of glucose tolerance was caused solely by insufficient insulin. Knock down of let-7 was sufficient to lead to increased insulin secretion and recovery of insulin sensitivity(36). Although being comprehensive and insightful, previous studies do not provide the information onlet-7in specific tissues and the function of let-7 in pancreatic 1-cells is incomplete and remains to be determined.

1.4 Methods to silence/block miRNA family function

miRNAs generally exist as multiple family members and display significant functional redundancy. Therefore, it is important to silence specific miRNA and reveal its biological function and its underlying regulatory networks. Currently, there are three approaches used for loss of miRNA function: genetic knockouts, miRNA sponges and antisense oligonucleotides(40). The traditional methods for gene functions by gene knockout are not very effective in studies of miRNA because of the small sizes and multiplicity of miRNA genes distributed over the
intergenic regions (41). miRNA sponges are transgenes that express in cells and generate the transcripts containing multiple miRNA binding sites. The promoter-driven sponges are allowed to contain a fluorescence reporter gene for identification and selection (42). Another approach in miRNA loss-of-function studies is to introduce chemically synthesized antisense oligonucleotides, or antimiRs. The antimiRs are presumed to anneal to and block the mature miRNA through sequence complementarities (43).

In this study, we construct the short tandem target mimic (STTM) to block the functions of miRNA. STTM is a powerful technology complementary to miRNA sponges. STTM is an artificial non-coding RNA consists of two miRNA binding sites and a spacer linker (Figure 1.4). The binding sites for a miRNA are perfectly complementary to their targets with a bulge at the central three nucleotides in order to prevent miRNA mediated cleavage and degradation. The spacer is 48-88 nucleotides long with a weak hairpin secondary structure (41, 44). Effectiveness of STTM has been confirmed by STTM-165/166 in plant (45) and STTM-30 in pancreatic 1-cells (44).

![Figure 1.4 Structure of STTM RNA transcripts.](image)

**Figure 1.4 Structure of STTM RNA transcripts.** A 48-88 nt spacer with the stem loop structure is flanked by two small RNA binding sites. The bulge
introduced into binding sites stabilizes the interactions with miRNA and prevents itself from being cleaved. Adapted from (44).
2. **Objective**

In this study, the objective was to identify the biological functions of let-7 microRNA in pancreatic 1-cells by blocking expression of endogenous let-7 using STTM methods. The hypothesis is that changes of let-7 in pancreatic 1-cells can result in significant differences in cell proliferation, cell death as well as insulin secretion of 1-cells. The mainly objective of this work is to examine the target gene regulated by let-7 and investigate the molecular mechanisms underlying the biological alterations.
3. Materials and methods

Construction of STTM-let7

To block the expression of both let-7a and let-7b in pancreatic 1-cells, a STTM sequence complementary to let-7a (AACTATAAACCctaTACTACCTCA) and let-7b (AACCACACAACCctaTACTACCTC) with an 88 nt spacer in between (total 138 bp) was synthesized and cloned into the pEGP-miR vector as described (44) (Figure 3.1).

Figure 3.1 Schematic representations of pEGP-miR cloning and expression vector for STTM-let7a/b and pEGP-miR Null control vector. (A) pEGP-miR (4972bp). The two sites for cloning of STTM in Human 1-globin intron are BamHI and Nhel. Positions indicating EF-1a promoter, GFP-Puro fusion protein, SV40 Polyadenylation signal and the human 1-globin intron are shown in the diagram. (B) pEGP-miR Null (4.7kb). The pEGP-miR Null control vector cannot be digested with BamHI due to secondary structure. (C) The sequences of STTM-let7a/b cloned into pEGP-miR vector. Two restriction enzyme recognition sites (BamHI and Nhel) are shown.

Cell culture and transfection
The pancreatic 1-cell line MIN6 was cultured in DMEM/high glucose (Thermo Scientific, Waltham, MA) with 15% fetal bovine serum (FBS) (Atlanta biologicals, Flowery Branch, GA), 100U/ml penicillin and 100 U/ml streptomycin. Cells were maintained in a 5% CO$_2$ humidified atmosphere incubator (Thermo Scientific, Waltham, MA) at 37 C. The culture medium was replaced every 48h. For transfection, MIN6 cells were trypsinized and pelleted at 700 rpm for 3 min. Cell pellets were resuspended in 100 µl Nucleofector buffer (4D Nucleofector X kit L) (Lonza, Basel, Switzerland). 10 µg of plasmids for pEGP-STTM-let7 and pEGP-Null control vector were used for transfection according to the protocol. 3 days after transfection, MIN6 cells were treated with cytokines mix (10ng/ml, IL-11, IFN-y and TNF-a) for 12-24h.

Isolation of pancreatic islets and virus infection

Islets were isolated from 10- to 16-week wild type mouse using collagenase digestion and cultured in RPMI medium. After incubating for 24h, islets were infected with recombinant adenovirus as previously described (46) and plated in 60 mm Petri-dish. Islets were collected 2 days after infection and perform western blot as described below.

Luciferase reporter assay

To evaluate the silencing efficiency of STTM-let7, let-7a complementary sequence or mutant was subcloned into the pRLTK vector (Promega, Madison, WI). For the luciferase reporter assay, pRLTK reporter constructs (2 µg) were electrorporated into MIN6 cells (10$^6$) with pEGP-STTM-let7 or pEGP-Null control vector using Amaxa (Lonza). The plasmid PGL-3 containing firefly luciferase (2µg) was co-transfected together to normalize for transfection efficiency. Luciferase activity was measured with a dual-luciferase reporter assay kit (Promega) two days after transfection.

RNA isolation and real-time PCR

Total RNA from MIN6 or islets was extracted using Trizol reagent (Roche, Basel, Switzerland) or miRNasy Mini Kit (QIAGEN, Venlo, Netherlands) and reverse-transcribed using TaqMan MicroRNA Reverse Transcription Kit (Applied
Quantitative RT-PCR was performed using the SYBR Green Master Mix and StepOnePlus Real-Time PCR System (Applied Biosystems). miRNA levels were normalized by the relative expression of U6.

**Western blot analysis**

MIN6 cells were cultured and treated as described above and lysed with lysis buffer containing 50 mM HEPES, 150 mM NaCl, 1 mM EDTA, 1% Triton, 0.2% SDS, phosphatase inhibitor and proteinase inhibitor (1 ml, Sigma, St. Louis, MO). Protein concentration was determined using PierceBCA Protein Assay Kit (Thermo Scientific) and 150µg total protein samples were separated by SDS-PAGE. Immunoblotting was performed with antibodies purchased from Cell Signaling (Beverly, MA) diluted according to the protocols.

**Cell proliferation assay**

DNA synthesis was measured using cell proliferation ELISA kit (Roche). MIN6 cells were transfected as described. At 48h after transfection, cells were collected and seeded in a 96-well plate to the density of 4x10^4 per well. BrdU were added to culture medium to a final concentration of 10µM. After BrdU labeling for 12-24h, the cells were processed according to the protocol.

**Cell Apoptosis**

At 48h after transfection, cells were collected and seeded in a 96-well plate. After incubating in the plate for 24h, cytokines mix was added to culture medium to the final concentration of 10ng/ml. According to the cell death detection ELISA (Roche) kit, anti-histone biotin and anti-DNA-POD were added to each well to measure the levels of cell death.

**Glucose Stimulated Insulin secretion**

As described above, MIN6 cells were seeded in 35 mm dish and incubated for 3 days followed by 12-24h treatment of cytokines mix. Afterwards, cells were pre-incubated for 2h in Krebs-Ringer bicarbonate HEPES buffer (KRB) and then cultured in KRB containing 25mM glucose for 1h. After the incubation period, supernatants were collected for stimulated insulin secretion and cell pellets were
processed for total insulin content. Insulin levels were measured using Mouse Insulin ELISA Kit (Mercodia, Uppsala, Sweden) and normalized to DNA concentration of the cell pellets.

**Statistical analysis**

For comparisons, statistical significance was evaluated using a two-tailed Student t test. A p value of less than 0.05 was considered significant.
4. Results

4.1 STTM-let7 reduces expression of let-7 in β-cells

The STTM-let7a/b constructed was synthesized and ligated with pEGR-miR vector that digested using BamHI and Nhel as described above. To investigate the expression of STTM-let7 and inhibition of endogenous let-7, pEGR-Null control vector and pEGR-STTM-let7 were transfected into MIN6 cells respectively. Cells were collected for Northern blot to detect the expression level of let-7a. Endogenous let-7a was down-regulated in cells transfected with STTM-let7 compared with control group (Figure 4.1 A). In order to further determine the silencing efficiency of STTM-let7, let-7a or mutant let-7 recognition sites were cloned into 3'UTR of pRLTK luciferase vector and luciferase reporter assays were performed. Co-transfection with STTM-let7 was capable of decreasing luciferase activity of the wild type constructs, whereas the reporter with mutant let-7 sites was not affected (Figure 4.1 B).

Figure 4.1 STTM-let7 reduces the expression of let-7. A: Northern blot analysis of total RNA from MIN6 cells transfected with pEGR-STTM-let7 or pEGR-Null control vectors with $^{32}$P-labeled let-7a probe (left). Let-7 level was normalized to U6 as fold change (right). B: the pRLTK-let7 luciferase reporter construct was co-transfected into MIN6 cells along with pEGR-STTM-let7 or pEGR-Null control, respectively. The luciferase activities were normalized by the co-transfected pGL3 firefly luciferase activity. **, P<0.01

4.2 Let-7 suppress cell death by targeting caspase-3
To understand the biological effects caused by the decrease of let-7 in 1-cells, we examined the expression level of let-7 targets by western blot. As expected, expression of caspase-3, which was reported to be one oflet-7 targets(19), was elevated when endogenous let-7 was inhibited by STTM (Figure 4.2 A, B) in MIN6 cells and islets. Caspase-3 belongs to the group of effector caspase family, which are located downstream of cell death pathway and can be activated through intrinsic pathway (Mitochondria) or extrinsic pathway (ligand-receptor binding)(17). Activation of Caspase-3 requires proteolytic cleavage of itself into 17kDa and 12kDa subunits. The intrinsic apoptotic pathway is mainly mediated by Bcl-2 family members that center at mitochondria. Bax and Bcl-2, two antagonistic Bcl-2 family proteins regulating apoptosis, were all increased by STTM-let7 independent of cytokines induced extrinsic cell death pathway. The elevation of Bcl-2 family by inhibition of let-7 suggested let-7 might play critical roles on regulating cell death.

The effect of let-7 on cell death was confirmed through cell death assay in MIN6 cells. In the presence of cytokines treatment, inhibition of let-7 by introducing STTM-let7 significantly increased cell death, which was correlated to the increased Caspase-3 expression (Figure 4.2 C). Under normal growth conditions, MIN6 cells displayed a low level of cell death rate and inhibition of let-7 had no effect.
Figure 4.2 Inhibition of let-7 by STTM-let7 induces cell death by targeting caspase-3. A: STTM-let7 increased the expression of caspase-3 and Bax. MIN6 cells were transfected with STTM-let7 or control vector. After 48 h, cells were treated with cytokine (10 µg/ml) for 16 h and the expression of caspase-3, cleaved caspase-3, Bcl-2 and Bax were measured by Western blot. B: STTM-let7 increased the expression of caspase-3 and Bax in isolated islets infected with AD-STTM-let7 virus. C: STTM-let-7 Increased cytokine-induced cell death in MIN6 cells. *, P<0.05.

4.3 Let-7 promotes 1-cell proliferation

To assess the contribution of let-7 to cell proliferation, BrdU incorporation was performed in MIN6 cells after transfection. After incubating for 24h, STTM-let7 was capable of decreasing BrdU signal and inhibiting cell proliferation (Figure 4.3 A). For the cells treated with cytokines, there was no difference in BrdU signal because of the intensive cell death rate. Western blot analysis indicated that IRS-
2 and mTOR, two factors involved in insulin signaling and regulate 1-cell function, were both down-regulated by STTM-let7 (Figure 4.3 B). However, the expression of Akt and phosphor-Akt was not changed significantly. Akt2 was predicted containing the miRNA recognition sites of let-7 and it was phosphorylated by mTOR complex 2 (mTORC2) at Serine 473 (47). Activation of Akt2 subsequently phosphorylates mTOR at Serine 2448 and results in activation of mTOR complex 1 (mTORC1). On the other hand, one of the insulin transcription factors, MafA, was not affected by STTM-let7 as expected for the reason of lacking let-7 recognition sites. However, STTM-let7 had no significant effects on the expression of mTOR, IRS-2 and Akt in islets with AD-STTM-let7 (data not shown).

**Figure 4.3 Inhibition of let-7 by STTM-let7 inhibit 1-cell proliferation by down-regulating insulin signaling.** A: MIN6 cells were transfected with STTM-let7 or control vector. After 48 h, cell proliferation was measure by BrdU incorporation after incubated with cytokine (10 μg/ml) for 16 h. B: Western blot validated that STTM-let7 decreased the expression of IRS-2, p-Akt, p-mTOR and mTOR. The expression of MafA had no significant effect.

### 4.4 Let-7 positively regulates glucose-stimulated insulin secretion

Glucose stimulated insulin secretion assay of MIN6 cells was performed to investigate the effects of let-7 on 1-cell function. Increased glucose level is
capable of inducing insulin secretion by causing production of ATP and elevation of cellular Ca\(^{2+}\), and finally leads to the release of synthesized insulin from secretory vesicles. Consistent with the repressed insulin signaling, STTM-let7 inhibited glucose-stimulated insulin secretion as well as total insulin content (Figure 4.4 A, B). Given the previous studies about insulin exocytosis was associated with insulin signaling (15), IRS-1 and PI3 kinase act as regulators controlling intracellular Ca\(^{2+}\) released from endoplasmic reticulum. As a result, the inhibition of insulin signaling in 1-cells also contributes to the reduction of insulin secretion.

Figure 4.4 Inhibition of let-7 by STTM-let7 decreased insulin secretion in pancreatic beta cells. MIN6 cells were transfected with STTM-let7 or control vector. After 48 h, the secreted insulin in the medium (A) and insulin content (B) were quantified using mouse insulin ELISA and normalized to total DNA. The presented data are the average of three independent experiments ± S.D.
5. Discussion

In this study, we observed that the expression of let-7 family was successfully blocked by the newly developed STTM technology. Inhibition of let-7 by STTM-let7 reduced 1-cell proliferation and induced 1-cell death through Caspase-3 mediated apoptosis pathway. Moreover, reduced let-7 expression impaired insulin signaling and decreased glucose-stimulated insulin secretion. Taken together, let-7 promotes 1-cell proliferation and protects against cytokine-induced apoptosis. Let-7 also plays a key role in regulating insulin secretion by activating insulin signaling.

Although let-7 was revealed to regulate apoptotic cell death by negatively regulating Caspase-3, whether other factors involved in cell death signaling are affected is less known. In this study, we show that let-7 inhibits two factors controlling initiation of cell death, Bax and Bcl-2. The elevation of Bax and Bcl-2 can be observed in MIN6 cells transfected with STTM-let7 with or without cytokines treatments, indicating let-7 regulates cells death independent of cytokine-induced extrinsic pathway. Cell death ELISA assay indicates that up-regulation of Caspase-3 by STTM-let7 sufficiently lead to a significant increase in cell death after cytokines inducement. Caspase-3 exists as an inactive form that is triggered by extracellular or intracellular stimulations through activation of initiator caspases (Caspase 8, 9 and 10) (17). Although increased Bax/Bcl-2 ratio was observed associated with increased Caspase-3 and apoptosis (48), the other caspases (caspase-8 or -9) downstream from Bax/Bcl-2 were not affected in the studies where caspase-3 was inhibited by overexpression of let-7(19). It could be the reason that Bax and Bcl-2 are up-regulated at the same extent and Bax/Bcl-2 ratio remains the same without affecting the downstream cell signaling. Whether let-7 inhibits the expression of Bax and Bcl-2 directly is unknown, however, let-7plays a role in regulating the center of cell death, Caspase-3.

Based on previous reports that let-7 can regulate glucose metabolism through Lin28/let-7 axis (30), let-7 controls glucose homeostasis and affects insulin sensitivity in muscles and liver by suppressing insulin signaling pathways. Insulin receptor (INSR), IRS-2, insulin-like growth factor 1 receptor (IGF1R), mTOR and Akt2 were confirmed to be regulated by let-7 through luciferase reporter assay (30,36). However, the previous studies primarily focused on insulin action on
insulin-target tissues rather than insulin secretion. Here, let-7 was revealed to promote insulin secretion and insulin signaling in 1-cells. IRS-2, which capable of promoting 1-cell growth and function (13), could play critical roles in let-7 mediated activation of insulin signaling. Decreased insulin secretion observed in cells transfected with STTM-let7 can be caused by inactivation of insulin signaling or impaired cell proliferation. Frost and Olson (36) have reported the decreased insulin secretion in let-7 globally knock-out mice. Theoretically, absent of let-7 in pancreas should promote insulin signaling and result in enhanced 1-cell function. However, a low level of insulin secretion in transgenic mice was observed, suggesting the potential effects of let-7 on maintaining glucose homeostasis. In our study, the inhibition of IRS-2 and mTOR in the presence of STTM-let7 indicates the reversal of let-7 effects on the mRNA of these genes. In contrast, the other let-7 target, Akt2 was not changed. The other isoform of Akt, Akt1 was ever revealed to regulate cell cycle by regulating Cyclin D1 and cyclin-dependent kinase-4 (49). Both of Akt1 and Akt2 were activated through mTORC2-mediated phosphorylation at Serine 473(47) although only Akt2 was predicted to be regulated by let-7. Absence of the alteration in Akt and p-Akt could be caused by the slight change of Akt expression and the antibody targeting total Akt content. The unaffected phosphor-Akt also suggests phosphorylation of Akt was not influenced by the decreased mTOR in mTOR Complex 2. Although an intact mTOR Complex requires phosphorylation of mTOR, recent data renders that mTOR is phosphorylated diversely when associated with mTORC1 and mTORC2 (50). The specific phosphorylation of mTOR in mTORC1 is primarily on Serine 2448 whereas mTOR in mTORC2 is phosphorylated on Serine 2481. It is mTORC2 that phosphorylate Akt1 and Akt2 at Serine 473. mTORC1 locates downstream from Akt and is activated through insulin signaling. Phospho-mTOR at Serine 2448 was detected and revealed to be repressed because of STTM-let7 (Figure 4.3 B). Unaltered phosphor-Akt provides the evidence that decrease of phosphor-mTOR only results from the low expression of mTOR, regardless of the activation of phosphor-Akt. Indeed, Ser-2448 phosphorylation of mTOR was reported to be regulated by p70S6 kinase downstream from mTOR (51).

Other studies demonstrate that the effect of let-7a on target mRNA is reversed in immune-stimulated cells (52) and translation of HMGA2 is activated by let-7 in
the cells arrested in G₀ phase of cell cycle (53). Bhattacharyya, et al discovered that the reversal of miRNA repression was mediated by mRNA processing bodies (P bodies) in response to different stress conditions (54). However, it is not confirmed whether the activation of insulin signaling by let-7 results from stress-induced reversal of mRNA repression. And in islets consist of alpha cells and 1-cells, IRS-2 and mTOR wasn't changed significantly by STTM-let7, suggesting let-7 may play different roles in different types of cell.

The decrease of insulin secretion caused by down-regulation of let-7 in MIN6 cells can be a combination of inhibited 1-cell growth and deactivation of insulin signaling. Changes of total insulin content suggest the let-7 promotes insulin expression by regulating cell proliferation. And essentially, insulin signaling also influences the Ca²⁺ level and directly affect the exocytosis of insulin vesicles (15). As mentioned above, increase of Ca⁺ was revealed partially dependent on IRS-1 and PI3 kinase (55). Recent data also indicates various intracellular factors involved in cell signaling contribute greatly to GSIS in compared with glucose itself (56). From the previously studies regarding the effects of insulin molecules on insulin secretion (15), it can be inferred there is a complex feedback mechanism that helps 1-cells auto-regulate their functions through insulin signaling.
6. Future work

In this study, the effects of let-7 on pancreatic 1-cells were mainly focused on MIN6 cell line. And the molecular mechanisms underlying the biological changes were not fully understood. In future work, STTM-let7 can be used for repression of let-7 in mice islets. Insulin promoter region will be used driving the expression of STTM in the 1-cell of mouse pancreas. Whether let-7 regulates 1-cell function can be determined in vivo by examining glucose metabolism in the transgenic mice. Globally inhibition of let-7 in mouse had been shown associated with increased insulin sensitivity and enhanced glucose tolerance (36). For the mouse specifically inhibit let-7 in 1-cells, glucose tolerance tests (GTT) will be performed to confirm the response of 1-cell to glucose stimulation.

The action of let-7 in other tissue cells should be examined for comparison with let-7 in 1-cells. Insulin signaling pathways in different tissues may be responsible for various functions and thus they are mediated in different manners. Another strategy resolving the question is to silence one of the genes involved in insulin signaling. Let-7 mediated 1-cell function could base on either a combination of multiple let-7 target genes or a few critical genes. In addition, the activation time of insulin signaling presumably leads to diverse effects on insulin secretion in response to glucose (15).

Although well known as a tumor suppressor, let-7 was recently reported to regulate cell death by targeting Fas (57) and Caspase-3 (19). Our work further revealed let-7-mediated apoptosis was associated with Bax/Bcl-2 ratio in mitochondria. It is essential to examine the miRNA involved cell death signaling in other types of cell. Overexpression of let-7 in those cells will provide more information about let-7 effects on cell growth and cell death.

Finally, 1-cell mass can be measured in isolated islets from the transgenic mice specifically silence let-7 in pancreas. Considering the low proliferation rate and apoptosis rate of 1-cells, whether absence of let-7 will result in a significant difference is unknown. Since let-7 is responsible for regulating glucose homeostasis and insulin sensitivity, dysfunction of let-7 in 1-cells may also contribute to the development of type 2 diabetes.

7. Conclusion
Short tandem target mimic (STTM) is capable of inhibiting the expression of endogenous microRNA let-7 in pancreatic 1-cells. Let-7 miRNA family plays significant roles on the normal function of 1-cells. Through STTM approach, let-7 was shown to protect 1-cells from cytokine induced cell death by suppressing Bax-Caspase-3 apoptotic cascade. Moreover, for the first time let-7 was revealed to regulate insulin signaling by controlling the expression of several factors responsible for 1-cell proliferation and insulin secretion. IRS-2 and mTOR were both found to be promoted by let-7. The study of let-7 in pancreatic 1-cells suggests the function of insulin signaling pathways in insulin-producing cells and provides potential targets that may contribute to therapeutic treatments of type 2 diabetes.
8. References


Ser2481 is a Marker for Intact mTOR Signaling Complex 2. Cancer Research 69, 1821-1827


