Stable Knockdown of Adenosine Kinase by Lentiviral Anti-ADK miR-shRNAs in Wharton’s Jelly Stem Cells

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Abstract

Objective: In this study, we describe an efficient approach for stable knockdown of adenosine kinase (ADK) using lentiviral system, in an astrocytoma cell line and in human Wharton’s jelly mesenchymal stem cells (hWJMSCs). These sources of stem cells besides having multigene differentiation potential and immunomodulatory activities, are easily available in unlimited numbers, do not raise ethical concerns and are attractive for gene manipulation and cell-based gene therapy.

Materials and Methods: In this experimental study, we targeted adenosine kinase mRNA at 3’ and performing coding sequences using eight miR-based expressing cassettes of anti-ADK short hairpin RNA (shRNAs). First, these cassettes with scrambled control sequences were cloned into expressing lentiviral pGIPZ vector. Quantitative real time-polymerase chain reaction (qRT-PCR) was used to screen multi-cassettes anti-ADK miR-shRNAs in stably transduced U-251 MG cell line and measuring ADK gene expression at mRNA level. Extracted WJMSCs were characterized using flow cytometry for expressing mesenchymal specific marker (CD44+) and lack of expression of hematopoietic lineage marker (CD45-). Then, the lentiviral vector that expressed the most efficient anti-ADK miR-shRNA, was employed to stably transduce WJMSCs.

Results: Transfection of anti-ADK miR-shRNAs in HEK293T cells using CaPO4 method showed high efficiency. We successfully transduced U-251 cell line by recombinant lentivirus and screened eight cassettes of anti-ADK miR-shRNAs in stably transduced U-251 MG cell line by qRT-PCR. RNAi-mediated down-regulation of ADK by lentiviral system indicated up to 95% down-regulation of ADK. Following lentiviral transduction of WJMSCs with anti-ADK miR-shRNA expression cassette, we also implicated, down-regulation of ADK up to 95% by qRT-PCR and confirmed it by western blot analysis at the protein level.

Conclusion: Our findings indicate efficient usage of shRNA cassette for ADK knockdown. Engineered WJMSCs with genome editing methods like CRISPR/cas9 or more safe viral systems such as adeno-associated vectors (AAV) might be an attractive source in cell-based gene therapy and may have therapeutic potential for epilepsy.

Keywords: Adenosine Kinase, Gene Knockdown Techniques, Lentivirus, RNA Interference, Wharton’s Jelly

Introduction

Previous molecular studies indicated that up-regulation of adenosine kinase (ADK), a key enzyme in the metabolism of adenosine, is one of the most important processes involved in astrogliosis (1, 2). RNA interference (RNAi) or post-transcriptional gene silencing (PTGS) is an interesting molecular tool for gene knockdown. Knock down of ADK increases intracellular adenine and results in extracellular adenosine augmentation. Adenosine has known protective effects on the central nervous system (3, 4). ADK gene could be targeted by RNAi in human cells which is an effective way to produce adenosine-releasing cells (5, 6). Adenosine augmentation exhibits a paracrine therapeutic effect and has potential for therapeutic applications in neurological diseases like refractory epilepsy (7).

Among children, the highest incidence of epilepsy is seen at ages less than five years old. Therefore, finding a new source of cells with therapeutic applications is highly required (8). Wharton’s jelly stem cells (WJMSCs) are an alternative for bone marrow mesenchymal stem cell (BMSCs). They are multipotent cells which are easily isolated in unlimited numbers with long-term ex vivo proliferation and immunomodulatory properties (9). WJMSCs are obtained from discarded human umbilical cord, with no ethical concern (10). These cells express specific MSCs markers like CD44 and are negative for CD45 hematopoietic lineage marker (11, 12). Being easily accessible, makes WJMSCs an alternative...
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In the present study, we used anti-ADK microRNA (miR) in a shRNA lentiviral system (miR-shRNA) for ex vivo gene therapy in U-251 MG cell line. We screened eight cassettes of miR-shRNAs that target human ADK gene. In order to screen and select the most efficient anti-ADK miR-shRNA, astrocytoma cell line was employed. Human U-251 MG cell line highly expresses ADK gene. Pseudo lentiviruses of eight anti-ADK miR-shRNAs were used for transducing of astrocytoma cell lines. The most efficient anti-ADK miR-shRNA for knockdown of ADK was selected by quantitative real time-polymerase chain reaction (qRT-PCR) analysis of established cells. Furthermore, human WJMSCs was confirmed by western blot analysis as well as qRT-PCR after transduction using the most efficient anti-ADK miR-shRNA lentiviral vector.

Materials and Methods

In this experimental study, human U-251 MG cell line (Sigma-Aldrich, USA) was cultured with Dulbecco’s Modified Eagle’s Medium (DMEM, Gibco-BRL, Japan) and 10% fetal bovine serum (FBS, Gibco, USA). This cell line highly expresses ADK gene. The third passage of these cells was used for screening the anti-ADK miR-shRNAs to knockdown ADK gene. All the experiments including animal works were approved by TUMS Ethics committee No. 93-01-87-25045-109011 and were performed based on the committee guideline.

Lentiviral constructs for the expression of anti-ADK miR-shRNAs

The eight different pre-miRNA sequences and a randomized scrambled control (SC) sequence were purchased (GE Healthcare). All miR-shRNA cassettes were cloned into the pGIPZ lentiviral vector, which contained a TurboGFP green fluorescence protein (tGFP) as a reporter gene, internal ribosome entry site (IRES) and a puromycin resistance gene; thus, it allowed co-expression of the respective miR-shRNA with tGFP and selection of stably transduced cells with puromycin. Expression of tGFP, puromycin and miR-shRNAs were under Cytomegalovirus (CMV) constant promoter. All genes were expressed as a single mRNA. At first, mRNA was processed in nuclear for producing premature miR-shRNA and bicistronic GFP-puromycin mRNA. pGIPZ lentiviral expression vector harbored internal long terminal repeats (LTRs) zeocin selection marker for selection of correct intact vector during bacterial propagation.

Production of recombinant pseudo lentiviruses

Recombinant lentiviruses were produced according to the Prof. Trono lab protocol with some modifications (13, 14). Briefly for all 8 miR-shRNAs and the positive control vector, 1×10⁶ HEK 293T cells (Invitrogen, USA) were cultured in a 10-cm² plate in DMEM medium supplemented with 10% FBS one day prior to transfection. Two hours before transfection, the medium was replaced with fresh medium. The transfection mixture contained 21 μg of pGIPZ-miR-sh/SC, 15 μg of pCMV-dR8.2, 10.5 μg of pMD2, 33 μl of TE 1X, 105 μl of 2.5 M CaCl₂, and 1064 μl of 2X HEPES-buffered saline (HeBS) finally reaching a volume of 2100 μl using buffered water.

All the components were mixed, and HeBS 2X was added when the solution was being vortexed vigorously. The final volume of transfection mixture used for each 10-cm² plate, was 2100 μl. HEK 293T cells were incubated with transfection solution in 37°C for 14 hours. The transfection medium was replaced with fresh medium 14 hours after transfection and cells were assayed for GFP expression using a fluorescent microscope (Labomed, USA). GFP expression indicated transfection efficiency. To estimate the transfection rate, five fields were randomly observed under the fluorescent microscope. Supernatant of cells containing recombinant viruses was collected at three time points (24, 48 and 72 hours after transfection) after 14 hours post-transfection. Next, collected supernatants were centrifuged at 180 g and filtered through a 0.25 μm filter before concentration and titration. For concentration of the supernatants, we used PEG 6000 and 9000 g centrifuge. The lentiviral titers from the 9 different miR-shRNA constructs due to tGFP gene in our vectors were determined using flow cytometry method. The determination of the lentiviral titer allowed us to estimate the multiplicity of infection (MOI) and thus to reduce the infectious activity of the viral stocks.

Determination of lentivirus titration

Since the stocks of vector carry GFP transgene that can be easily monitored by flow cytometer, we used this method for titration of lentivectors. We used Prof. Trono lab protocol for lentivirus titration (15). For this purpose, 293T cells were cultured in DMEM medium supplemented with 10% FBS in a 12-well plate with 1×10⁵ cells in each well. Then, the medium was removed and cells were transduced in 500 μl of fresh DMEM with serial dilutions of the vector that corresponded to the final amount of 1, 10⁻¹, 10⁻², 10⁻³ and 10⁻⁴ μl of the vector. After 24 hours, the medium was removed and one ml of fresh medium was added to each well. Then, 72 hours after transduction, cells were processed for Fluorescence Activated Cell Sorting (FACS) analysis. The following formula was used for calculating titer by flow cytometry:

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\text{Titer HEK 293T transducing U/ml}=\frac{\text{Number of target cells (counted on day 1)×(% of GFP-positive cells/100)\times MOI\times Volume\ of\ supernatant}}{\text{Volume\ of\ transfection\ mixture}}
\]

Ideally, the cells should present 2-20% GFP to confirm that cells that received multiple copies of the virus, were not counted. The flow cytometer apparatus did not have enough sensitivity for determination of GFP <1%.

Genetic engineering of U-251 MG cell line

After three passages, U-251 MG was cultured in six-well cell culture plates at a density of 2×10⁵ cell /well /2 ml. Before adding fresh recombinant viruses, U-251 MG cell line was washed with PBS. Recombinant viruses in culture medium with an MOI of 5 were used for transduction of cells in each well. To increase the rate of transduction, the spinfection method was used. After infection, the plates were incubated at
37°C and the medium was changed 24 hours later. Transduced cells were assayed for GFP expression with a fluorescent microscope (LaboMed, USA), 72 hours after transduction. GFP expression indirectly indicated the expression of miR-shRNA and transduction efficiency. To select stably transduced cells, from day 4, the medium was replaced with fresh medium containing 2 µg/ml puromycin for 5-7 days.

Quantitative analysis of ADK knockdown by real-time analysis in U-251 MG cells

A QuantiFast SYBR Green RT-PCR Kit (Qiagen, USA) was employed to determine and monitor ADK expression after knockdown of this gene by recombinant lentiviral constructs that express anti-ADK miR-shRNAs. After puromycin selection, 2×10⁶ cells from stable U-251 MG cell line was used for RNA extraction and cDNA synthesis (Qiagen, kit, USA). According to the manufacturer protocols, cDNA was used for standard real-time PCR. Specific primer pairs were used for ADK and TBP (TATA Sequence-Binding Protein). The expression of ADK mRNA was evaluated in the lentiviral-engineered U-251 MG cells. For improving the reliability of relative RT-PCR, TBP was used as a reference gene.

Isolation and expansion of human Wharton’s jelly mesenchymal stem cells

Human Wharton’s jelly tissues were obtained from newborn umbilical cord in accordance with bioethics agreement following obtaining the consent from its mother at Taleghani Hospital, Tehran, Iran. Mucoid connective tissue or Wharton’s jelly were separated from blood vessels (two arteries and one vein of the umbilical cord) then, washed twice with PBS containing penicillin, streptomycin and amphotericin. The cord was rinsed with PBS and isolated from amniotic membrane. After separation of the matrix from cord vessels, the jelly matrix was cut into small pieces and transferred into culture dishes supplemented with DME/F12 medium, 10% FBS and antibiotics. Two weeks later, the tissues were discarded and the isolated growing cells (WJMSCs) were fed with the same medium. The cells were grown to 60% confluence and passaged by trypsinization. The third passage of WJMSCs was used for characterization.

Flow cytometry

A number of 1×10⁵ cultured WJMSCs were washed, fixed and incubated for 15 minutes at 4°C with a 1:9 dilution of normal goat serum in phosphate buffered saline (PBS, Sigma-Aldrich, USA). Then, cells were incubated for 1 hour with FITC-conjugated antibodies (CD44 and CD45) for labelling. Cells washed with 2% FBS in PBS were used for analyzing with FACS Calibur apparatus (Becton Dickenson, USA). The control population was stained with isotype-matched antibodies (FITC-conjugated and PE-conjugated mouse IgG monoclonal isotype standards) and confirmed by positive fluorescence of the linbal samples. For each sample, at least 1×10⁶ events were recorded and analyzed by WinMDI software (USA).

Quantitative analysis of ADK knockdown by selected anti-ADK miR-shRNA in Wharton’s jelly mesenchymal stem cells

A QuantiFast SYBR Green RT-PCR Kit (Qiagen, Germany) was employed to monitor knockdown of ADK after transduction of WJMSCs by the most efficient recombinant lentiviral construct that expressed anti-ADK miR-shRNA. After puromycin selection with 2 µg/ml for 5-7 days, 1×10⁶ cells from WJMSCs were used for RNA extraction and cDNA synthesis (Qiagen, Germany). In the second step, cDNA was used for standard real-time PCR. Specific primer pairs were used for ADK and TBP. The expression of ADK mRNA was evaluated in the lentiviral-engineered WJMSCs cells.

Preparation of total protein extracts

Stable transduced umbilical cord mesenchymal stem cells (WJMSCs) were lysed using ReadyPrep™ mammalian cell lysis reagent (Bio-Rad) comprising complete protease inhibitor cocktail. Cellular proteins were prepared according to manufacturer’s protein extraction protocol. The samples were stored at -80°C until western blot analysis. The protein content of sample lysate was measured using Bradford’s method (16).

Western blot analysis

An equal amount of proteins was loaded on 12% sodium dodecyl sulfate-polyacrylamide gel electrophoresis (SDS-PAGE) and transferred to polyvinylidene fluoride membranes. Blots were blocked using non-fat dry milk for 1 hour. After blocking, membranes were probed with ADK primary antibody (Santa Cruz, 1/1000). Then-anti-rabbit IgG antibody conjugated with horseradish peroxidase (Santa Cruz, 1/8000) was used for 1 hour at room temperature. Subsequently, the blots were reprobed with a β-actin antibody (1:2000, Abcam, ab8227, USA). Then, the blots were treated with Electro Chemi Luminescence reagents (Amersham Biosciences, UK). For quantification of protein intensities, western blot bands were visualized on radiograph film and evaluated by Image J software. Human umbilical cord tissue was used in accordance with the Declarations of Tehran University of Medical Science and Stem Cell Research Center Committee after obtaining written informed consent from the mother.

Results

Transfection efficiency of lentiviral anti-ADK miR-shRNAs vectors in packaging 293T cell line

All of nine transfer vectors (eight anti-ADK miR-shRNAs and scrambled control) with helper vectors (pCMV-dR8.2 and pMD2) were separately co-transfected into HEK293T cell line using CaCl₂ protocol. The transfection efficiency, as evaluated based on GFP marker under the fluorescent microscope, was more than 90-95% for eight anti-ADK miR-shRNAs (Fig.1) and scrambled control.

Titration of lentivirus by FACS

The titer of viral particles was approximately 1.5-2×10⁷ U/ml. Serial dilutions of the vector that corresponded to the final amount of 10⁻¹⁻¹⁰ μl of the vector, were used for calculating the titration because the GFP+ cells in other
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serial dilutions were more than 20% or under 1%.

Transduction of U-251 MG cell line

As U-251 MG cell line highly expresses ADK gene, we chose it for screening anti-ADK miR-shRNAs (Fig.2). Nine recombinant pseudo lentiviruses (eight anti-ADK miR-shRNAs and scrambled control) were separately used for transduction of U-251 MG cell line. GFP reporter gene showed high efficiency of transduction and expression of anti-ADK miR-shRNAs and scrambled pseudo lentiviruses in U-251 MG cells (Fig.3). Selection of GFP+ cells, 72 hours after transduction, was done by using puromycin (2 µg/ml) (Fig.4).

Fig.1: Transfection efficiency of lentiviral anti-ADK miR-shRNAs vectors in packaging 293T cell line. A-H. HEK 293T cells were transfected with anti-ADK miR-shRNAs and I. Scrambled control with high efficiency (90-95%). The rate of GFP expression evaluated undera fluorescent, microscope confirmed the high efficiency of transfection and expression of anti-ADK miR-shRNAs.

Fig.2: Morphological characteristics of U-251 MG cells. A. These cells are shown under a phase-contrast microscope with low confluence and B. High confluence.
Fig. 3: Lentiviral Transduction efficiency in U-251 MG cell line. High rate of tGFP expression was seen under a fluorescent microscope 72 hours after transduction with anti-ADK miR-shRNAs lentiviral vectors. A-H. Transduction with sh1-sh8 and I. Transduction of cells with scrambled control.

Fig. 4: Antibiotic selection of transduced U-251 MG cell line. The stable cell line was produced by puromycin selection after 7 days. A, B. Depict the second and third day after using puromycin (2 µg/ml), C and D. Show the 5th and 7th day of selection.
Screening of anti-ADK miR-shRNAs in U-251 MG cells by semi-quantitative real-time polymerase chain reaction

For selection of the most efficient anti-ADK miR-shRNAs for ADK knockdown, quantitative real-time PCR was employed. Expression of ADK gene in each of target groups (U-251 MG cell line transduced with 8 anti-ADK miR-shRNAs) was measured in proportion to control group (U-251 MG cell line transduced with scrambled control). For improving the reliability of relative RT-PCR, TBP was used as a reference gene. Analysis by quantitative real-time PCR was done by REST2009 software (Fig.5A). Data showed that anti-ADK miR-shRNAs, except for sh8, could knockdown ADK more than 60% and sh4 and sh7 were the most efficient anti-ADK miR-shRNAs with 86 and 95% knockdown of ADK, respectively (Fig.5B).

Morphological characteristics and mesenchymal stem cell markers expression of human Wharton’s jelly cells

Umbilical cord matrix tissue was cultured by an explant method. A few days later, WJMSCs migrated away from tissues. After two weeks, fibroblast-like cells appeared in culture dishes. The cells grew fast and rapidly covered the surface. Flow cytometric analyses indicated that the cultured WJMSCs do not express hematopoietic marker CD45; however, they expressed mesenchymal stem cell marker CD44 (Fig.6A).

Genetic engineering of WJMSCs by anti-ADK miR-sh7

Analysis of real-time data showed that Anti-ADK miR-sh7 is the most efficient (up 95%) anti-ADK miR-shRNA for knockdown of this gene. For this reason, we used pseudo lentiviruses of this miR-shRNA for transduction of WJMSCs. The high efficiency of transduction was seen under the fluorescent microscope. After selection with puromycin (1.5 µg/ml) in the culture medium (Fig.6B), semi-quantitative real-time PCR was employed for measuring ADK expression at mRNA level. Data also showed knockdown of ADK gene in WJMSCs up to 95% as well as down-regulation in the U-251 MG cell line.

Confirm of ADK knockdown in WJMSCs by western blot analysis

After selection of transduced WJMSCs using anti-ADK miR-sh7 lentiviral vector, cell lysates were used for performing western blot. Cell lysates from transduced WJMSCs with scrambled control viruses, cell lysates from WJMSCs and HepG2 as a control were also employed. To quantify and normalizing of ADK immunoreactivity, β-actin antibody were used. Analysis of data did not show any reduction of ADK in WJMSCs, WJMSCs-miR-shSC, and HepG2; however, down-regulation of ADK was observed after transducing cells with the human specific anti-ADK miR-sh7 (Fig.6C).
Fig.6: Knock down of ADK in human Wharton’s jelly cells. A. Morphological characteristics and mesenchymal stem cell markers expression of human Wharton’s jelly cells. WJMSCs showed fibroblast-like phenotype under a phase-contrast microscope (a, b). Flow cytometric analyses indicated that the cultured WJMSCs significantly express mesenchymal stem cell marker CD44 (c). These cells were almost negative for hematopoietic marker CD45 (d).

B. Genetic engineering of WJMSCs by anti-ADK miR-sh7. WJMSCs observed under phase-contrast microscope (b). Stable transduced WJMSCs with anti-ADK miR-sh7 under fluorescent microscope (a). Puromycin- selected WJMSCs transduced with anti-ADK miR-shSC (1.5 μg/ml) (c).

C. Western blot analysis confirmed ADK knockdown in WJMSCs. Western blot analysis was performed on cell lysates from WJMSCs, WJMSC- miR-sh7, WJMSC-miR-shSC, and HepG2. ADK staining (top) and β-actin staining (bottom). WJMSC- miR-sh7 showed the most marked reduction of ADK expression at protein level.

Discussion

In adult brain, ADK is a key enzyme in astrocytes that regulates adenosine level by converting adenosine to 5′-adenosine monophosphate (AMP) and subsequently generates ATP (17). When trauma or epilepsy happens, adenosine, as a modulator of inflammation increases and induces proliferation of astrocytes via A2ARs, leading to astrocyte activation and ADK overexpression. ADK expression results in adenosine decline. In an acute injury or intractable epilepsy, proliferation of astrocytes (i.e. astrogliosis), is the pathological hallmark of the disease and at the molecular level, adenosine deficiency is the consequence of ADK overexpression induced by over-activation of astrocytes (18). So, ADK down-regulation and increasing adenosine release are among the strategies used for the treatment of some neurological disorders like epilepsy. Although inhibition of ADK with chemical and small molecule drugs can suppress seizures but systemic long-term use of this therapeutic approach may increase the risk of side effects like brain hemorrhage (19). Post-transcriptional gene silencing is a molecular tool that has opened new windows in recent years. Knockdown of ADK by viral vectors that express anti-ADK miRNAs.
can increase adenosine and results in suppressing seizures via a paracrine effect (20). Epilepsy is a chronic disease so permanent release of adenosine is needed. In this therapeutic approach, the combination of cell therapy with gene therapy is the key to the riddle.

Cell therapy provides long-lasting focal delivery of anti-epileptic molecules like adenosine and prevents systemic pharmacological side effects. Cell therapy also has the potential to restore cells that are destroyed by neurological conditions, especially in refractory epilepsy. In cell therapy approaches, safer and more controllable gene delivery of inhibitory neurotransmitters or other therapeutic compounds is possible. Chemical mutations for engineering cells to release therapeutic agents like adenosine, were induced in 2001. In that study, fibroblasts that release adenosine were generated by induction a deficiency in ADK and adenosine deaminase (ADA) gene (21). Transplantation of these cells implicated nearly complete protection against seizures up to 24 days but lost their anti-epileptic effects after that as the viability of grafting fibroblast cells decreased. Stem cells are more attractive sources to be used for solving this problem. Stem cells in addition to having higher viability, can be differentiated into multi-lineage cells such as neuro-progenitor cells to integrate into damaged neural networks that are created during epilepsy or other neurological disorders, they can slow seizure development and they have anti-epileptogenesis effects by releasing inhibitory substances.

Different sources of stem cells (e.g. adult stem cells, fetal stem cells, embryonic stem cells and iPS cells) have been used to evaluate treatment of animal model of epilepsy (22-25). Therapeutic application of ADK gene knockout in mouse embryonic stem cells and ADK gene knockdown in bone marrow mesenchymal stem cells have been investigated by Li et al. (26). These studies implicated reduction of seizures in a mouse model after transplantation of hMSCs and the results were promising to be used for treatment of epilepsy. However, ethical concerns about embryonic stem cells and technical issues related to bone marrow mesenchymal stem cells, have made WJMSCs as an attractive source for cell therapy and gene delivery systems.

WJMScs could be obtained from the human umbilical cord that is normally discarded after the birth. These cells are actually a by-product of childbirth. WJMScs have the potential to differentiate into ectodermal and mesodermal derived cells (27, 28). In addition to the easy access to WJMScs and their potential of differentiation, these cells have anti-inflammatory and immunomodulatory properties. Some studies have shown that xenograft transplantation of WJMScs are not rejected in an animal model of human disease without immune-suppression (29). Studies suggested that production of cytokines and growth factors by WJMScs, results in their anti-inflammatory and neuro-protective activities (30). In the present study, we improved therapeutic benefit of WJMScs by knocking down ADK gene and producing adenosine releasing WJMScs to suppress seizures in families with status epilepticus and we stored these cells for possible future applications. WJMScs are an attractive alternative source for cell-based gene therapy as they are i. Easily accessible stem cell sources without ethical and technical concerns, ii. Accessible in unlimited numbers and, iii. Amenable to genetic modification, and iv. They possess therapeutic potential. Based on these considerations, we selected these cells for engineering with lentiviral anti-ADK miR-shRNA expressing systems.

miRNA-mediated downregulation of ADK was previously reported by Li et al. (26). They used five anti-ADK miRNA expression cassettes and showed downregulation (up to 85%) of ADK in human MSCs (hMSCs). In the present study, we considered human miR 30 for designing stem-loop and flanking sequences of anti-ADK miR-shRNAs to increase the efficiency of RNAi-based gene therapy. We used eight anti-ADK miR-shRNAs that target ADK at different sense and antisense sequences but at common stem-loop and flanking sequences.

By developing this system, we succeeded to diminish ADK expression up to 95% in astrocytoma cell line and in WJMScs. ADK mRNA has 4 variants (isoform a, b, c and d). These eight anti-ADK miR-shRNAs can target all transcript variants of ADK. Antisense of sh6 and sh8 target ADK mRNA at 3’ outside of coding sequence (CDs) while other shRNAs (sh1, sh2, sh3, sh4, sh5 and sh7) target ADK mRNA at 3’ including CDs. Using human miR30 for designing shRNAs caused 60-95% knockdown of ADK (except for sh8 which resulted in 23% downregulation). Anti-ADK miR-sh4 and Anti-ADK miR-sh7 were the most efficient cassettes with up to 86 and 95% downregulation of ADK, respectively. These results show that the usage of miR-based shRNAs is an efficient method in knocking down ADK. Based on these results, in our future research, we work on using engineered WJMSc for the survey of the therapeutic potential of ADK down-regulation and adenosine delivery in an animal model.

Conclusion

Results show that lentiviral system expressing anti-ADK miR-shRNAs that was used in this study is a promising tool for ADK knockdown in all transcript variants. Manipulated WJMScs by Adeno, AAV, non-integrated lentiviral, CRISPR/Cas9 and RNA transfer instead of lentivirus, might be a suitable source in cell-based and ex vivo gene therapy of epilepsy and could be evaluated for ADK knockdown, in vivo.

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Author’s Contributions

M.H.Gh.; Supervised the project and conceived the study and were in charge of overall direction and planning. A.F.; Contributed to conception, design and revising the manuscript. H.E.; Worked out almost all of the technical details and wrote the manuscript. M.S.; Helped in the primary design of the experiments. A.A.; Helped in animal surgeries and stem cell isolation. F.K.; Analysed the data of EEG. Sh.B.A.; Contributed to the western blot experiment. All authors read and approved the final study.

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