Pancreatic cancer (PaCa) is currently the third most frequent cause of cancer-related death in the United States [1]. Pancreatic ductal adenocarcinoma (PDAC), the major histological subtype comprising 90% of all PaCa, displays local invasion and metastasis during early stages of the disease via developing intrinsic resistance to most therapeutics, contributing to its notoriously poor prognosis with 1–5% of 5-year survival rates (~6 months of median survival duration with currently available therapies) [2–4].

Natural killer (NK) cells constitute a significant component of the innate immune system which contributes particularly to the anti-tumor immune response [5, 6]. NK cells recognize virus-infected and transformed malignant cells and then kill these target cells without prior sensitization [6, 7]. NK cells have various surface receptors (NK cells are CD3-negative, CD16- and/or CD56-positive) and cytotoxic granules. NK cells recognize tumor cells and are thought to be good candidates for adaptive immunotherapy against cancer [8]. Recently, they gained attention for their spontaneous cytotoxicity against a broad range of malignancies and for their potential to become a “multi-purpose” anti-cancer agent and studies are focused on isolating and expanding NK cells abundantly [1, 6, 8]. NK cells exist in the peripheral blood at a very low concentration and after isolation there is a need to expand them in culture in order to reach a sufficient concentration for clinical applications. However, prolonged culture leads to exhaustion of NK cells and they become ineffective in killing target cells and die within a few days in clinical applications. Therefore, in recent years there have been great interests in attempts to generate NK cells from more abundant cell sources. Until today, NK cells have been differentiated in vitro from different cell sources such as embryonic stem cells (ESCs) or hematopoietic stem cells (HSCs) [6]. NK cells have been also directly isolated from umbilical cord blood or peripheral blood cells [6, 9]. Mesenchymal stem cells (MSCs) are multipotent stem cells with promising potential for cancer research [10–12]. Recently, these cells draw attention because of their importance for tissue microenvironment and their ability of suppressing immune system [11]. They can be obtained from many tissues such as bone marrow and adipose tissue, etc. and are able to be proliferated or differentiated suitably in in vitro conditions [13, 14]. Usually adipose tissue can be obtained easily and abundantly by a surgical procedure, which is not severely invasive and provides a rich source of MSCs [15, 16]. In addition, adipose derived MSCs (AD-MSCs) have less ethical problems when compared with other sources.

MicroRNAs (miRNAs) are short (~22 nt) noncoding regulatory RNAs that function as post-transcriptional...
miR150 is a microRNA that can act as a repressor of protein-coding target messenger RNAs and regulate translational repression or degradation [17, 18]. miR150 is expressed in various immune cells such as mature NK and innate NK T cells, B and T cells and is up-regulated during cellular maturation [19]. Particularly miR150 is responsible for the development and activation of NK cells and regulates the production of IFNγ by NK cells [19].

The aim of this study is to differentiate AD-MSCs to NK-like cells by using miRNA transfection in addition to classical differentiation protocol and then to investigate the effect of these cells on human PaCa cells in vitro.

**MATERIALS AND METHODS**

**Cell culture.** AD-MSCs purchased from ATCC were cultured in MesenPRO (Thermo Fisher Scientific, MA, USA) medium. Human pancreatic duct endothelial cells (hTERT HPDE) (a kind gift of MD Anderson Cancer Center, TX, USA) were cultured in keratinocyte serum free medium (Thermo Fisher Scientific, MA, USA) supplemented with 10% fetal bovine serum (FBS), 2 mM L-glutamine, 10 ng/ml of endothelial growth factor and 1% penicillin/streptomycin (P/S) (BI, Cromwell, CT, USA). Human PaCa cell line (Panc1) was obtained from MD Anderson Cancer Center, TX, USA and cultured in DMEM medium supplemented with 10% FBS, 2 mM L-glutamine and 1% P/S. AD-MSC derived NK-like cells were cultured in RPMI-1640 medium supplemented with 10% FBS, 12.5 ng/ml IL-2, 5 ng/ml IL-15 (Sigma, USA) and cultured in PaCa cells and cancer-related genes such as KRas, TP53, CDKN2A, SMAD4, TGFBR2, ARID1A by RT-PCR in PaCa cells was analyzed (Roche, LightCycler 480 Instrument, Basel, Switzerland). The primers used are given in Table.

**Reverse transcription-polymerase chain reaction (RT-PCR).** Total RNA was isolated from miR150-transfected and non-transfected AD-MSC-NK cells and PANC1 cells before and after co-culturing using RNeasy Mini Kit (Qiagen, Hilden, Germany). Expression of NK cells-specific genes such as GZMB, KIR2DL2, CD16, CD56, NKG2D and Nkp46 in NK-like cells and cancer-related genes such as KRas, TP53, CDKN2A, SMAD4, TGFBR2, ARID1A by RT-PCR in PaCa cells was analyzed (Roche, LightCycler 480 Instrument, Basel, Switzerland). The primers used are given in Table.

**Table. Primers used for RT-PCR**

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
</tr>
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<tbody>
<tr>
<td>GZMB</td>
<td>5'-GAAACGCTACTAAGCT-3'</td>
<td>5'-CCATCATGCTAAGAGG-3'</td>
</tr>
<tr>
<td>KIR2DL2</td>
<td>5'-AGGGTTTCTGCTGCT-3'</td>
<td>5'-GGGATCCCCAGGTG-3'</td>
</tr>
<tr>
<td>CD16</td>
<td>5'-CCCCAGTGGAGG-3'</td>
<td>5'-GGGATCCCCAGGTG-3'</td>
</tr>
<tr>
<td>CD56</td>
<td>5'-TGTGGTTTTTGGGAACTCC-3'</td>
<td>5'-CCGCGTCAGGATGTTG-3'</td>
</tr>
<tr>
<td>NKG2D</td>
<td>5'-TGGACTGAGCTGATGCTTTG-3'</td>
<td>5'-GGATATTGATGTTG-3'</td>
</tr>
<tr>
<td>Nkp46</td>
<td>5'-GGCTGTTGCTGAG-3'</td>
<td>5'-GATGAGTGCCGCTGAT-3'</td>
</tr>
<tr>
<td>KRAS</td>
<td>5'-AAGGGCCTCTGGAAATGACTG-3'</td>
<td>5'-GAACTGATGTTAGGC-3'</td>
</tr>
<tr>
<td>TP53</td>
<td>5'-AAGGCAGCTGCCACCC-3'</td>
<td>5'-GGGTGCCTCCGTTGGA-3'</td>
</tr>
<tr>
<td>CDK21A1</td>
<td>5'-AGAGAGAGGGGCTGCTGCT-3'</td>
<td>5'-GAGGCGCCGCCCTTCTCTT-3'</td>
</tr>
<tr>
<td>SMAD4</td>
<td>5'-TTGAGTTGGGCGCTGGA-3'</td>
<td>5'-CAGGTTGATAACCTGCTTT-3'</td>
</tr>
<tr>
<td>TGFBR2</td>
<td>5'-TGGGGAGAGGGCCGCT-3'</td>
<td>5'-GGCTGCGTCTGCACGGAGG-3'</td>
</tr>
<tr>
<td>ARID1A</td>
<td>5'-CTTACCATCCTAG-3'</td>
<td>5'-GGTACCGCACTCATACTTT-3'</td>
</tr>
<tr>
<td>Beta Actin</td>
<td>5'-GACACCAACCTTCTA-3'</td>
<td>5'-TGTTGCTGTAACCACTTCTACG-3'</td>
</tr>
</tbody>
</table>

**Immunohistochemical analysis.** The cells were fixed with 4% paraformaldehyde for 30 min and were washed two times with phosphate buffer saline (PBS). They were then incubated 10 min with 0.1% Triton X-100 solution on ice for permeabilization and 3% hydrogen peroxide (H2O2) was applied for 5 min after washing with PBS. They were then treated with blocking solution (Invitrogen, CA, USA) for an hour at room temperature and were incubated with primary antibody anti-hNK22D (RD Systems, MN, USA) overnight at 4 °C. The samples were washed with PBS and incubated with biotinylated rabbit anti-mouse secondary antibody (Invitrogen, CA, USA) for 30 min. After washing with PBS streptavidin-hydrogen peroxidase (Invitrogen, CA, USA) was added for 30 min. In order to develop the immunohistochemical reaction, diamobenidine (DAB) (ScyTek, UT, USA) was applied for 5 min. After washing with PBS, slides were stained with Mayer’s hematoxylin and mounted with mounting medium. The intensity of immunolabeling was evaluated by the two investigators in different times with light microscopy (BX40, Olympus, Tokyo, Japan). The immunoreactivities were considered as negative (-), weak (+), moderate (++) and strong (+++).
(Sigma, MO, USA) in a ratio of 1:2 (NK:PANC1). After incubation for 1 week, the number of dead cancer cells was determined via TUNEL assay.

**TUNEL assay.** The *in situ* apoptosis detection kit (ApopTag Plus Peroxidase In Situ Apoptosis Detection Kit, Millipore, MA, USA) was used. The cells were fixed with 4% paraformaldehyde for 30 min and were washed two times with PBS. They were then incubated 10 min with 0.1% Triton X-100 solution on ice for permeabilization and endogenous peroxidase activity was inhibited with 3% H$_2$O$_2$. The cells were then incubated with equilibration buffer for 10–15 s and TdT-enzyme in a humidified atmosphere at 37 °C for 60 min. They were subsequently put into pre-warmed working strength stop/wash buffer at room temperature for 10 min and incubated with anti-streptavidin-peroxidase for 45 min. Each step was separated by careful washing in PBS. Staining was performed with DAB and counterstaining was performed in Mayer’s hematoxylin.

**Statistical analysis.** Statistical analysis of the data was performed using the Mann–Whitney *U* test using the SPSS 16 (Statistical Package for the Social Sciences) program. The statistical significance was evaluated, considering *p* < 0.05 level as statistically significant.

## RESULTS

**Hematopoietic induction and NK-like cell differentiation.** AD-MSCs in culture were observed as fusiform cells adherent to the plastic surface (Fig. 1, *a*). When cultured in hematopoietic induction medium for a week, AD-MSCs detached from plastic surface and became suspended after forming sphere clusters (Fig. 1, *b*). Hematopoietically induced-AD-MSCs were then cultured in NK differentiation medium for 4 weeks in order to obtain NK-like cells and these cells were called AD-MSC-NK cells (Fig. 1, *c*).

**Characterization of cultured cells was performed immunohistochemically.** Distributions of CD90 and CD314 (NKG2D) were assessed following 4 weeks of culturing of AD-MSCs and AD-MSC-NK cells (Fig. 2). CD90 immunoreactivity was very weakly positive (Fig. 2, *e*) while NKG2D (CD314) immunoreactivity was strongly positive in AD-MSC-NK cells (Fig. 2, *f*) at the 30th day of culture. In control slides, negative staining for both CD90 and CD314 was detected (Fig. 2, *a*, *b,*).

Following miR150 transfection of NK cells differentiated from AD-MSC, NKG2D immunoreactivity was found to be strongly positive for miR150-AD-MSC-NK cells in comparison to non-transfected AD-MSC-NK cells (Fig. 3).

![Fig. 1. Characterization of AD-MSC: *a* — AD-MSC culture; *b* — hematopoietic induction; *c* — AD-MSC-NK cell culture](image)

![Fig. 2. Characterization of AD-MSC (*a*, *c*, *e*) and AD-MSC-NK (*b*, *d*, *f*) cells. Control staining (*a*, *b*), CD90 (*c*, *d*), CD314 (NK-G2D) (*e*, *f*)](image)

![Fig. 3. Immunoreactivity of CD314 (NKG2D) in AD-MSC-NK cells non-transfected (*a*) and miR150-transfected (*b*)](image)

Expression of NK cell specific genes such as GZMB, KIR2DL2, CD16, CD56, NKG2D and NKp46 significantly increased in AD-MSC-NK cells (*p* < 0.05) when
compared to undifferentiated AD-MSC as the control group (Fig. 4) and following miR150 transfection, the expression of these genes was found to be even greater in the miR150-transfected AD-MSC-NK cells (miR150-AD-MSC-NK) than the non-transfected AD-MSC-NK cells (AD-MSC-NK) (Fig. 5).

**Induction of apoptosis in PANC1 cells after co-culturing with AD-MSC-NK cells.** The TUNEL method based on detection of DNA strand breaks to identify apoptotic cells was used. According to our TUNEL assay results, AD-MSC-NK and miR150-AD-MSC-NK were both able to kill PANC1 cancer cells (Fig. 6). TUNEL-positive cells were not detected in the control staining of PANC1 cells, whereas 11% of TUNEL-positive cells were observed in PANC1 cells after co-culturing with NK cells (Fig. 6, b). After miR150 transfection of NK cells, the percentage of TUNEL-positive cells was 8.8% in miR150-NK-PANC1 co-culture (Fig. 6, d) but only 3.0% in miR150-NK-hTERT co-culture (Fig. 6, f). Only 4% of TUNEL-positive cells were found in PANC1-AD-MSC co-culture (Fig. 6, h).

After PANC1 cells were co-cultured with AD-MSC-NK cells, the expression of KRAS oncogene significantly decreased ($p < 0.05$), while the expression of tumor suppressor genes such as $P53$, $CDKN2A$, $TGFBR2$, and $ARID1A$ significantly increased ($p < 0.05$) in comparison with that in PANC1 cells before co-culturing (Fig. 7). After miR150 transfection, expression of tumor suppressor genes further increased while the oncogene expression was further reduced in PANC1 cells co-cultured with the miR150-transfected AD-MSC-NK cells in comparison to co-culture of PANC1 cells with non-transfected AD-MSC-NK cells (Fig. 8).

**DISCUSSION**

PaCa, which is frequently seen in the elderly, is an aggressive progressing cancer with 5-year
specific surface marker CD314 (NKG2D) were evaluated as well as genes such as GZMB, KIR2DL2, CD16, CD56, and ARID1A, or the patient were used [22].

In our study, the expression of NK cell-specific genes such as GZMB, KIR2DL2, CD16, CD56, NKG2D, and Nkp46 and immunoreactivity of NK cell-specific surface marker CD314 (NKG2D) were evaluated in NK-like cells before and after performing nanoparticle mediated miR150 transfection in AD-MSC-NK cells. Both miR150-transfected and non-transfected NK cells were co-cultured with PaCa cells (PANC1) in order to evaluate their efficiency in killing cancer cells and our results showed that miR150-transfected NK cells and non-transfected NK cells have similar apoptotic activities. However, RT-PCR results indicated that the decrease in gene expressions of oncogenes and the increase in the expression of tumor suppressor genes of PaCa were more prominent in co-culturing PaCa cells with miR150-transfected NK cells in comparison to that in co-culturing PaCa cells with non-transfected NK cells.

The results of the immunohistochemical analysis showed that CD314/NKG2D staining was strongly positive in miR150-NK and AD-MSC-NK cells compared to AD-MSCs. RT-PCR analysis indicated that GZMB, KIR2DL2, CD16, CD56, NKG2D, and Nkp46 genes expressed in NK cells were found to be more potent in the miR150-transfected cells.

According to TUNEL results, both miR150-transfected AD-MSC-NK and non-transfected AD-MSC-NK-like cells were similarly effective in inducing apoptotic death of PaCa cells. While cells gone to apoptosis can be analyzed by TUNEL assay, the pre-apoptotic cells cannot be analyzed by this assay so we performed Annexin V in our continuing study and determined 30% of pre-apoptotic cells in Annexin V assay (data not given).

It should also be noted that there is another pathway leading to cell death named pyroptosis, which is a proinflammatory apoptosis pathway. It will also be valuable to evaluate the pyroptosis pathway in PANC1-NK cells co-culturing via adding caspase-1 analysis to the procedures already used in our study. IL-18, an inflammatory cytokine associated with NK-like cells, is active in the pathway of pyroptosis, via activating caspase-1, which is not effective in the known apoptosis pathway but is effective in pyroptosis [23].

MSCs are derived from tissues that contain a rich source for the surrounding connective tissue. And thus AD-MSCs are promising immunomodulator stem cells which can be obtained easily and abundantly from several tissues, in particular by liposuction and lipectomy procedures [24, 25], and AD-MSCs do not carry any legal problems also the efficiency of NK cells can be increased via differentiation process.
mice [28]. In our study, NK-like cells differentiate from AD-MSCs were transfected with miR150 using nanoparticles for the first time. RT-PCR analyzes showed that in both non-transfected NK cells and miR150-transfected NK cells oncogene KRAS expression decreased, while expression of tumor suppressor genes (P53, CDKN2A, TGFB2, SMAD4 and ARID1A increased).

In a recent study, Joshi and colleagues investigated the effect of miRNAs responsible for regulating activation and function of NK cells on melanoma cancer stem cells. In their study, NK cells isolated from the peripheral blood and then transfected with anti-miR155 or anti-miR146 both showed antitumorigenic effect on melanoma cells and increased production of antitumorigenic factors released from NK cells such as INFγ and granzyme B [29]. In another study, Bezman and colleagues demonstrated that mice with a targeted deletion of miR150 have an impaired, cell lineage — intrinsic defect in their ability to generate mature NK cells while miR150 transgene promotes the development of NK cells, with more mature phenotype and also more responsive to activation indicating that miR150 transfection enhances the function of NK cells [19]. In accordance with these findings, in our study, it was shown that miR150 increased the expression of NK cell-specific genes as it was shown in Fig. 3 and 5.

The efficiency of NK cells derived from HCSs obtained from fresh or frozen cord blood samples was investigated in cancer cell lines using three different culture systems developed by Spanholtz et al. According to their data, the most effective NK cells were obtained from frozen cord blood HSCs and these cells can be used in cancer treatment [30]. A study by Luévano and his team supported these results. It has been reported that NK cells obtained from frozen cord blood HSCs were more effective [9].

Since the activity of NK cells in cancer patients diminishes or disappears with time, cytokines which have potential in activating NK cells such as IL-2, IL-12, IL-15, IL-18, IL-21 are systemically administered to the patients in order to induce endogenous NK cells [31]. However, in these treatment approaches high doses of these cytokines are administered to the patients to strengthen and enumerate NK cells, and it is known that this high dose cytokine treatment has systemic toxic effects [32]. In our study, while obtaining NK-like cells from AD-MSCs which has potential for allogeneic use, we used cytokines at concentrations such as 50 ng/ml SCF, 50 ng/ml Flt-3 ligand, 12.5 ng/ml IL-2, 20 ng/ml IL-7, 40 ng/ml IL-15 for 4 weeks in vitro during the differentiation protocol. In our study, when performing differentiation protocol in vitro we used IL-2 and IL-15 at the nanogram levels which would not lead to systemic toxicity in the future even we will continue to use cytokines at low levels during in vivo studies.

Since PaCa is one of the most aggressive cancers, our results are promising in improving immunotherapeutic approaches to cure malignancies. The results of this study are encouraging for further clinical studies in improvement of immunotherapeutic approaches for treatment.

ACKNOWLEDGEMENT

This study was supported by grant No. 214S650 from the Turkish Scientific and Technological Research Institute (TUBITAK).

REFERENCES


