Human adipose mesenchymal stem cell-derived exosomal-miRNAs are critical factors for inducing anti-proliferation signalling to A2780 and SKOV-3 ovarian cancer cells

Abu Musa Md Talimur Reza, Yun-Jung Choi, Hideyo Yasuda & Jin-Hoi Kim

An enigmatic question exists concerning the pro- or anti-cancer status of mesenchymal stem cells (MSCs). Despite growing interest, this question remains unanswered, and the debate became intensified with new evidences backing each side. Here, we showed that human adipose MSC (hAMSC)-derived conditioned medium (CM) exhibited inhibitory effects on A2780 human ovarian cancer cells by blocking the cell cycle, and activating mitochondria-mediated apoptosis signalling. Explicitly, we demonstrated that exosomes, an important biological component of hAMSC-CM, could restrain proliferation, wound-repair and colony formation ability of A2780 and SKOV-3 cancer cells. Furthermore, hAMSC-CM-derived exosomes induced apoptosis signalling by upregulating different pro-apoptotic signalling molecules, such as BAX, CASP9, and CASP3, as well as downregulating the anti-apoptotic protein BCL2. More specifically, cancer cells exhibited reduced viability following fresh or protease-digested exosome treatment; however, treatment with RNase-digested exosomes could not inhibit the proliferation of cancer cells. Additionally, sequencing of exosomal RNAs revealed a rich population of microRNAs (miRNAs), which exhibit anti-cancer activities by targeting different molecules associated with cancer survival. Our findings indicated that exosomal miRNAs are important players involved in the inhibitory influence of hAMSC-CM towards ovarian cancer cells. Therefore, we believe that these comprehensive results will provide advances concerning ovarian cancer research and treatment.

Different organs including ovaries are surrounded and supported by adipose fat-pad, which provide physical, as well as mechanical supports and play important roles during organogenesis, morphogenesis, disease-progression of respective organs. Being an important composite of adipose-stromal cells, adipose mesenchymal stem cells possibly have regulatory part in different cancers, such as ovarian cancer. However, relationships between mesenchymal stem cells (MSCs) and cancer cells are a mystery, owing to insufficient evidence concerning both the stimulatory and inhibitory roles of MSCs on cancer cells. While there is debate about the customary roles of MSCs, their involvement in cancer biology is undoubtedly clear. MSCs potentially support tumour development through immune suppression, epithelial-to-mesenchymal transition, angiogenesis, and serving as cancer stromal cells. In contrast, MSCs also suppress cancer by downregulating cancer survival-signalling pathways involving WNT/β-catenin and/or AKT. There is a need to investigate the mechanisms underlying the contradictory roles associated with MSCs in cancer biology. Cytokines and soluble factors secreted by MSCs have been thoroughly scrutinized, with most reports concluding that MSC-secreted cytokines and soluble factors exhibit stimulatory effects related to cancer progression.

Exosomes are types of membrane-bound micro-vesicles 30 nm to 200 nm in diameter, found in bio-fluids and contain many important components, including RNA, proteins, DNA, and lipids, and serve as efficient vehicles for cancer-stromal communication. Exosomes are secreted by all cells and, despite their ability to be incorporated...
into neighbouring cells, have been only marginally investigated. Specifically, cell-secreted microRNAs (miRNAs; 18–22 nucleotides) are predominantly carried by exosomes and have been studied in recent years for their roles in post-transcriptional regulation of gene expression through miRNA silencing. Therefore, understanding the functions of the MSC-derived secretome (particularly exosomes) in cancer is critical to elucidating the cross-talk between MSCs and cancer cell biology.

In this study, we hypothesized that human adipose-derived MSC (hAMSC)-secreted biological component (cytokines, miRNAs and others) might have important influence on the regulation of ovarian cancers. Hence, we investigated the influence of hAMSC-secreted molecules on different ovarian cancer cells. Our results showed that hAMSC-conditioned medium (hAMSC-CM)-derived exosomes treatment inhibited the proliferation and growth of A2780 and SKOV-3 ovarian cancer cells. More precisely, cancer cells exhibited reduced viability, wound healing, and colony formation following fresh or protease-digested exosome treatment; however, treatment with RNase-digested exosomes could not inhibit the proliferation of A2780 and SKOV-3 cancer cells. Furthermore, sequencing of exosomal RNAs revealed a rich population of miRNAs, with many reported to exhibit anti-cancer properties through targeting different cancer-survival pathways. Our findings indicated that exosomes (particularly exosomal miRNAs) may be one explanation for the anti-proliferative effects exhibited by hAMSC-CM, and that the relationship between MSCs and cancer could be partially explained by exosome-related activity. These results provided valuable insights into the diversity, enrichment, and function of all miRNAs derived from hAMSC-secreted exosomes.

**Results**

**hAMSC-CM treatment reduced proliferation of A2780 ovarian cancer cells.** Treatment with hAMSC-derived CM altered cell proliferation through enhanced oxidative stress and reduced mitochondrial membrane potential (MMP). During the course of determining optimal treatment parameters, we observed that supplementation with hAMSC-derived CM did not exhibit changes in pH of culture medium; however, as shown in Supplementary Fig. S1, cell-viability assays revealed that viability began to decrease following treatment with 20% CM, with optimum inhibition observed at 25% CM supplementation. As shown in Fig. 1a, significant decreases in cell viability were observed at 48 h (20% reduction) and 72 h (40% reduction) after treatment with 25% CM (p < 0.05 and p < 0.01, respectively) as compared with the control.

To evaluate the effects of cell apoptosis according to hAMSC-derived CM treatment, we examined the generation of reactive oxygen species (ROS) and MMP in CM-treated A2780 cells. As shown in Fig. 1b, 25% CM treatment resulted in an enhanced JC-1 monomer/aggregate ratio of 65% relative to control cells, as well as significant increases of ROS to 30% more than observed in control cells (p < 0.05) (Fig. 1c). Cell cycle analysis at 48 h following CM treatment revealed significant (p < 0.05) increases in the number of S-phase cells and decreases in G0/G1- and G2/M-phase cells (Fig. 1d). However, 72 h of CM treatment resulted in increases in both S-phase and G2/M-phase cells, and significant (p < 0.05) decreases in G0/G1-phase cells. These results indicated that hAMSC-CM restrained proliferation and blocked cell cycle of A2780 cancer cells.

**hAMSC-CM treatment activates the intrinsic apoptosis pathway in A2780 cells.** To determine the induction of cell death following hAMSC-derived CM treatment, first we performed tetramethylrhodamine (TMR)-red assay and measured the percentage of apoptotic cells. Fluorescence-activated cell sorting analysis revealed significant (p < 0.05) increases in the percentage of apoptotic cells following CM treatment as compared with control cells (Fig. 1e). Second, we decided to detect the intrinsic apoptosis pathway-related anti- and pro-apoptotic molecules expression in CM-treated A2780 cells. In CM-treated A2780 cells, expression of anti-apoptotic protein BCL2 was downregulated as compared with control cells, while, several pro-apoptotic molecules, including phosphorylated p53, BAX, CASP9, CASP3, and cytosolic Cytochrome-c expression were significantly upregulated or activated. Additionally, the BAX/BCL2 ratio and active-/pro-CASP3 ratios increased by several folds in CM-treated cells relative to controls (Fig. 1f). Supplementary Fig. S2 shows full-length of gel images. These findings indicated that hAMSC-CM treatment initiated activation of the intrinsic apoptosis pathway in A2780 cells.

**Identification and characterization of hAMSC-CM-derived exosomes, and incorporation of exosomes within A2780 cells.** Then, we investigated whether hAMSC-CM-derived exosomes were responsible for the induction of apoptosis observed on CM-treated A2780 cells. Figure 2a shows hAMSC-CM-derived exosomes detected by transmission electron microscopy (TEM). Exosomes appeared as spheres having a size distribution from 50 nm to 150 nm, although the majority appeared to be between 70 nm and 100 nm (Fig. 2b). Next, we extracted the total proteins from exosomes and visualized the results with Coomassie Blue Staining (Fig. 2c). As shown in Fig. 2d, CD63, which is an exosome protein marker, was detectable in the exosomes-derived lysate, verifying the status of these structures as exosomes. Finally, we inspected whether exosome can internalize into A2780 cancer cells. To visualize, exosomes were labeled using Exo-red stain kit, which signifies binding with exosomal RNAs. As expected, CM-derived exosomes shows extensive internalization into A2780 cancer cells (Fig. 2e), indicating that hAMSC-CM-derived exosomes were successfully absorbed by the cells and might participate in deciding the fate of A2780 cancer cells.

**hAMSC-CM-derived exosomes reduced cell viability, wound-repair capacity, and colony formation ability of A2780 & SKOV-3 ovarian cancer cells.** To determine the influence of hAMSC-CM-derived exosomes internalization on ovarian cancer cells, we treated A2780, SKOV-3 and CAOV-3 ovarian cancer cells using hAMSC-CM-derived exosomes and found that exosomes treatment had inhibitory influence on A2780 and SKOV-3 cells, while there was no remarkable change in CAOV-3 cells. We examined the wound-repair capacity (Fig. 2f) and colony formation ability (Fig. 2g) of A2780 cells. As shown in Fig. 2h,i and j, we observed significant
reductions in cell-plating efficiency, cell-survival fraction, and cell viability following exosome internalization as compared to control cells (p < 0.05, p < 0.01, and p < 0.01, respectively). To determine the effects of A2780 cell incubation with either CM, exosomes, or both, we observed that incubation with exosomes alone caused a 28% loss in cell viability as compared to control cells, whereas incubation with exosomes and CM resulted in a 33% decrease in cell viability (p < 0.01; Supplementary Fig. S3). Similarly, cells treated with CM alone decreased the wound-healing capacity of the cells relative to control cells, while cells that internalized exosomes or both exosomes and CM, exhibited far less efficiency in wound-healing (Supplementary Fig. S3). Control cells exhibited ~46% plating efficiency, whereas cells that treated with CM or exosomes or else both exosomes and CM, exhibited 30% to 35% plating efficiency (p < 0.05, p < 0.01, respectively; Supplementary Fig. S4). Similarly, cell survival are also decreased by 30% to 35% in cells incubated with CM or those that had internalized CM-derived exosomes (p < 0.05 and p < 0.01, respectively) relative to levels observed in control cells (Supplementary Fig. S4). As presented in Supplementary Fig. S5, treatment using hAMSC-CM-derived fresh exosomes restrained proliferation of SKOV-3 cells by around 20% similarly, protease-digested exosomes treatment also caused approximately 20% viability declination of SKOV-3 cells, compared to the control cells. In contrast, RNase-digested exosomes could not display inhibitory influence on SKOV-3 cells. These results indicated that hAMSC-CM-derived exosomes might be potential inhibitors against proliferation of A2780 and SKOV-3 ovarian cancer cells.

**A2780 cells responded dose- and time-dependently upon hAMSC-CM-derived exosomes treatment.** As shown in Supplementary Fig. S6, dose-dependent treatment using hAMSC-CM-derived exosomes and viability of A2780 ovarian cancer cells demonstrated a negative correlation, higher the concentration of exosomes, lower the cell viability. More clearly, gradual inhibitions of cell proliferation recorded with gradual increase of doses of hAMSC-CM-derived exosomes. Likewise, viability of A2780 cells significantly (p < 0.01) reduced at 24 h incubation with one U/mL hAMSC-CM-derived exosomes, and the maximum inhibition was recorded at 48 h of treatment.
hAMSC-CM-derived exosomal miRNAs inhibit A2780 cells proliferation. We then investigated whether exosomal protein or RNA was responsible for the anti-proliferative effects observed on A2780 cells. As shown in Fig. 2k, incubation of A2780 cells with protease-digested exosome lysates significantly reduced cell viability (p < 0.01) to a degree similar as that observed upon internalization of intact CM-derived exosomes. However, similar incubation with lysates from RNase-digested exosomes did not inhibit A2780 cell proliferation (Fig. 2l), but rather caused slight increases in cell viability. These findings suggested that exosomal RNA was responsible for inhibiting A2780 cell viability.

Next-generation sequencing (NGS) confirmed oncogene-related miRNAs in hAMSC-CM-derived exosomes. Presence of small RNA molecules in exosomes was confirmed by verifying the integrity of exosomal RNA (Fig. 3a), as well as by successful preparation of small RNA library (Fig. 3b) from hAMSC-CM-derived exosomal miRNAs. To assess whether miRNAs were present as precursors or fully processed transcripts, we examined length distributions of the miRNA reads and found that the majority of hAMSC-derived exosomal miRNA sequences ranged between 18 and 25 nucleotides in length (Supplementary Table S1).

Aiming to receive a quick impression about the quality of raw sequence data delivered by high throughput sequencing pipelines, quality control check-up was performed by FastQC (http://www.bioinformatics.babraham.ac.uk/fastqc).
with a quality filter Q-score > 50 (Fig. 3c). Total reads were detected between 20,633,027 and 22,988,601, with a clustered read count between 687,897 and 765,338 in all three samples where the detected mapped and unmapped sequences were ~2% and ~98%, respectively. We determined that a substantial proportion of the cellular RNA content from hAMSC-derived exosomes was small-RNA, with ~2% constituting known miRNA and 1.2% potential candidate miRNA. The data also indicated the presence of ribosomal RNA, transfer RNA, small nuclear RNA, and small nucleolar RNA (Fig. 3d).

We found 141 known miRNAs (Fig. 3e) and 25 novel potential miRNAs (Fig. 3f), which were common in all three replicates. The most abundantly expressed known miRNA was hsa-miR-4792, which had read counts of 361010, 381745, and 381918 in the three replicates, respectively (Table 1). However, role of hsa-miR-4792 in cancer have not reported in literature and, most probably, remains undefined. The second and third most abundant miRNAs were hsa-miR-320b and hsa-miR-320a, which belong to the hsa-miR-320 family known for its roles in different molecular-and oncogene-signalling pathways. Various members of the hsa-miR-127, hsa-miR-486, hsa-miR-423, hsa-miR-181, hsa-miR-1246, hsa-miR-26, and hsa-miR-378 families were also observed among the 20 most abundant miRNAs, with each widely investigated and reported as having roles in cancer-regulatory activity. Several other miRNAs which belong to top 20 known exosomal miRNAs, have not been extensively investigated for their oncogene regulatory properties; these miRNAs include hsa-miR-7704, hsa-miR-6087, hsa-miR-22-3p, hsa-miR-4466, hsa-miR-4532, hsa-miR-7641, hsa-miR-4448, hsa-miR-3960, and hsa-miR-3687 (Table 1).

The 141 known miRNAs belong to 104 different miRNA families, with 20 families containing two or more miRNAs found in all three replicates. Among them, eight miRNAs (Supplementary Table S1) belong to the hsa-let-7 family, which are recognized suppressor miRNAs in many types of cancer. As demonstrated in Supplementary Table S1, five miRNAs belong to the hsa-miR-1273 family, which does not have well-documented anti-/pro-cancer properties, while among detected four members of the hsa-miR-30 family, hsa-miR-30c is reportedly involved promoting the invasiveness of metastatic breast cancer cells. Five families (hsa-miR-320, hsa-miR-181, hsa-miR-29, hsa-miR-125, and hsa-miR-151) were each associated with three miRNAs found here. While members of the hsa-miR-320 family are reported as having anticancer properties, miR-151 family members are documented as exhibiting pro-cancer functions. Member(s) of the hsa-miR-181 family...
| miRNA           | Precursor    | Sequence                      | miRBase       | ENTREZGENE   | Sample 1 | Sample 2 | Sample 3 |
|----------------|--------------|-------------------------------|---------------|--------------|----------|----------|----------|
| hsa-miR-4792   | hsa-mir-4792 | cggugacgcucucgucgc           | miRBase:hsa- | ENTREZGENE:  | 3610     | 381745   | 381918   |
| hsa-miR-320b   | hsa-mir-320b 1,hsa-mir-320b-2 | aaagcgggggagagggcca          | miRBase:hsa-miR-320b | ENTREZGENE:  | 2628     | 2094     | 2046     |
| hsa-miR-320a   | hsa-mir-320a | aaacgcggggagagggcggca        | miRBase:hsa-miR-320a | ENTREZGENE:  | 1645     | 1311     | 1293     |
| hsa-miR-7704   | hsa-mir-7704 |ccccgcgccccgggccg             | miRBase:hsa-miR-7704 | ENTREZGENE:  | 1470     | 971      | 1145     |
| hsa-miR-127-3p | hsa-mir-127  |ugcgcugcuagucgcgucggc        | miRBase:hsa-miR-127-3p | ENTREZGENE:  | 943      | 836      | 807      |
| hsa-miR-6087   | hsa-mir-6087 |ggcucagugguauauggaaagccu     | miRBase:hsa-miR-6087 | ENTREZGENE:  | 783      | 831      | 885      |
| hsa-miR-22-3p  | hsa-mir-22   |aagcucggccagcagacgcgcu       | miRBase:hsa-miR-22-3p | ENTREZGENE:  | 772      | 638      | 671      |
| hsa-miR-4466   | hsa-mir-4466 |ggcggcgccggccccgggggg         | miRBase:hsa-miR-4466 | ENTREZGENE:  | 459      | 541      | 618      |
| hsa-miR-486-5p | hsa-mir-486-1,hsa-mir-486-2 |gucgcucgacucgucgccccggag     | miRBase:hsa-miR-486-5p | ENTREZGENE:  | 581      | 529      | 453      |
| hsa-miR-7641   | hsa-mir-7641-1,hsa-mir-7641-2 |ugcgcugcuagucgcu           | miRBase:hsa-mir-7641 | ENTREZGENE:  | 449      | 449      | 625      |
| hsa-miR-423-5p | hsa-mir-423  |ggcgcgcgcggaggggggggggggggggg | miRBase:hsa-mir-423-5p | ENTREZGENE:  | 518      | 490      | 428      |
| hsa-miR-3960   | hsa-mir-3960 |ggccgggacgggccccccuccgggcc   | miRBase:hsa-mir-3960 | ENTREZGENE:  | 328      | 273      | 273      |
| hsa-miR-1246   | hsa-mir-1246 |aaagcggcuacgcacgcggc        | miRBase:hsa-mir-1246 | ENTREZGENE:  | 226      | 224      | 130      |
| hsa-miR-26a-5p | hsa-mir-26a-1,hsa-mir-26a-2 |ccgacuccgaggcagccg        | miRBase:hsa-mir-26a-5p | ENTREZGENE:  | 195      | 143      | 155      |
| hsa-miR-378a-3p| hsa-mir-378a  |agccgacgcagagacgcgcu        | miRBase:hsa-mir-378a-3p | ENTREZGENE:  | 134      | 105      | 148      |

**Potential New-miRNAs**

| Potential New-miRNA | Mature Sequence | Precursor Sequence | Mature Length (nt) | Chr No. | Start | End | Precursor Length (nt) |
|---------------------|-----------------|--------------------|--------------------|---------|-------|-----|-----------------------|
| chr1:12001518504    | aagcgggaa        | aagcgggaa          | 18                 | chr12   | 12150| 12168| 18                    |
| chr2:1450164804     | ccgccggggcag     | ccgccggggcag       | 20                 | chr7    | 16951| 16951| 20                    |
| chr3:70092157       | ggccgacggg       | ggccgacggg         | 18                 | chr16   | 70002| 70021| 18                    |

**Expected Candidate-miRNAs**

| Expected Candidate-miRNA | Mature Sequence | Precursor Sequence | Mature Length (nt) | Chr No. | Start | End | Precursor Length (nt) |
|--------------------------|-----------------|--------------------|--------------------|---------|-------|-----|-----------------------|
| chr8:56821963-56822004  | puggcuccaaag     | puggcuccaaag       | 18                 | chr8    | 56821| 56822| 18                    |
| chr9:28678365-28678429  | pgggacgcgcgacgcg | pgggacgcgcgacgcg   | 19                 | chr6    | 28678| 28678| 19                    |
| chr10:28726148-28726212| pgggacgcgcgcggcgc| pgggacgcgcgcggcgc| 19                 | chr6    | 28726| 28726| 19                    |
| chr11:36065457-36065499| gcgcgcgcggcggggc| gcgcgcgcggcggggc  | 19                 | chr20   | 36065| 36065| 19                    |
| chr12:61801606-61801662| gcgcgcgcgcggcggc| gcgcgcgcgcggcggc  | 18                 | chr20   | 61801| 61801| 18                    |
and hsa-miR-29 families exhibit both anticancer and pro-cancer regulation under different clinical conditions, while members of hsa-miR-126, hsa-miR-129, hsa-miR-204, hsa-miR-656, hsa-miR-99 families are recognized as having anticancer properties under different clinical conditions. Families that contained at least one miRNA found in hAMSC-CM-derived exosomes reportedly exhibit antagonistic roles in numerous cancer types and include hsa-miR-486, hsa-miR-100, hsa-miR-101, hsa-miR-124, hsa-miR-125, hsa-miR-134, hsa-miR-138, hsa-miR-139, hsa-miR-143, hsa-miR-186, hsa-miR-193, hsa-miR-205, hsa-miR-222, hsa-miR-338, hsa-miR-339, hsa-miR-424, hsa-miR-127, hsa-miR-26, hsa-miR-103, hsa-miR-107, hsa-miR-128, hsa-miR-140, hsa-miR-144, hsa-miR-153, hsa-miR-192, hsa-miR-212, hsa-miR-218, hsa-miR-23, hsa-miR-25, hsa-miR-370, hsa-miR-375, hsa-miR-381, hsa-miR-410, hsa-miR-411, and hsa-miR-93. These NGS data revealed an enriched miRNA population in hAMSC-CM-derived exosomes that are likely involved in the regulation of different types of cancer.

**Functional enrichment analysis of exosomal miRNAs.** To broaden our search for potential miRNAs, gene ontology (GO) functional enrichment analysis was performed on 8195 genes predicted as targets of the 141 known miRNAs found in CM-derived exosomes (Supplementary Spreadsheet S1). GO analysis indicated that approximately 40% of the targeted genes were associated with cellular components, including 24% with organelles, 14% with the cell membrane, 10% with macromolecular complexes, 7% with extracellular regions, and 3% with the extracellular matrix. A small portion of molecules was also associated with cell junctions and synapses (Fig. 4a). Of the target genes associated with molecular functions, approximately 33% are involved as binding molecules, 29% with catalytic activity, and 5% to 10% of the remaining genes with enzyme regulation, nucleic acid binding/transcription-factor activity, receptor activity, structural activity, and transporter activity (Fig. 4b). GO terms associated with biological processes were more diversified, with 25% of the target molecules involved in metabolic processes, 21% in cellular processes, 13% in biological regulation, 8% in developmental processes, 6% in stimuli response, and 5% in multicellular organismal processes. Other targeted molecules were reportedly associated with biogenesis, apoptosis, reproduction, growth, biological adhesion, immune response, cell killing, and locomotion (Fig. 4c).

**In silico pathway analysis revealed cancer-related targets associated with hAMSC-CM-derived exosomal miRNAs.** According to Kyoto Encyclopaedia of Genes and Genomes (KEGG) pathway analysis, 173 pathways were detected as targets for the 141 miRNAs detected in CM-derived exosomes (Supplementary Table S2). Each of the pathways was targeted by multiple miRNAs, with five miRNAs targeting the lowest scoring pathway and 88 miRNAs targeting the top-scoring pathway, which was 'Pathways in cancer' (Fig. 4d). Molecules associated with many other signalling pathways were predicted as targets for exosomal miRNAs, including the cancer-survival pathways related to TP53, MAPK, WNT, JAK-STAT, PI3K-AKT, cAMP, mTOR, TGF-β, VEGF, and PPAR.

The target molecules predicted by BioCarta (http://www.genecarta.com/), including those related to cell death and BCL2-associated death promoter (BAD)-signalling, are presented in Supplementary Fig. S7. Twenty two molecules associated with 'Death-signalling pathway' were predicted as targets for the 141 miRNAs found in CM-derived exosomes and included BCL2, XIAP, IAP, CASP3, GAS2, APAF1, CASP7, CASP10, CYCS, CASP8, SPTAN1, LMNA, BIRC2, MAP3K14, FADD, NFKB1, TNFRSF10B, BID, CASP6, NFKBIA, RELA, and DFFB. Additionally, 20 molecules related to the BAD-signalling pathway included IGF1R, BCL2, IGF1, ADCY1, PIK3R1, MAPK1, KITLG, BCL2L1, YWHAA, PRKACB, RPS6KA1, KIT, PIK3CA, PRKAR2B, PRKAR2A, PIK3CG, AKT1, BAD, IL3, and PRKAR1A. These findings indicate that miRNAs found in CM-derived exosomes target molecules involved in both cancer-survival signalling and cell death pathways, suggesting potentially important roles in the regulation of cancer proliferation, progression, and metastasis.

**Validation of miRNA expression in hAMSC-CM-derived exosomes and host cells.** The expression of five selected miRNAs (hsa-miR-4792, hsa-miR-6087, hsa-miR-320a, hsa-miR-7704, and hsa-miR-181a-5p) was confirmed in both hAMSC-CM-derived exosomes and a host cell line by using quantitative reverse-transcription polymerase chain reaction (qRT-PCR). All five miRNAs were detected in both exosomes and the host cell line, with three miRNAs (hsa-miR-4792, hsa-miR-6087, and hsa-miR-181a-5p) downregulated and two miRNAs (hsa-miR-320a and hsa-miR-7704) upregulated in exosomes as compared with the host cell line (Fig. 4e). These data indicated that exosomal miRNA expression could reflect the miRNA expression of host cell line in terms of miRNA population diversity, but may not reflect the level of expression of different miRNAs.

**Validation of target molecules associated with cancer-specific pathways.** For target validation, each of the miRNAs (hsa-let-7a-5p, hsa-miR-181a-5p, hsa-miR-320a, hsa-miR-124-3p, and hsa-miR-26a-5p) was transfected into separate A2780 cells, respectively. Overexpression of specific miRNAs in each transfected cells were confirmed using qRT-PCR, with cells transfected with ‘mir Vana™’ miRNA Mimic negative Control.
considered as an experimental control. The mimic-miRNA transfected cells exhibited up to 500-folds greater levels of expression relative to levels observed in the negative-control cells (Fig. 5a–e). We then assessed changes in the expression of selected target genes in miRNAs-transfected cells versus those in negative-control cells, to verify miRNA targets. Our results revealed that several members of the cyclin-dependent kinase (CDK) family, including CDK2, CDK4, and CDK6 (Fig. 5f–h), were targeted by hsa-miR-124-3p. Furthermore, several cytokines and cytokine receptors genes were also targeted by different exosomal miRNAs; these genes include IGFR, TGFBR1, AR, ITGB1, and ITGB3, which were downregulated by hsa-let-7a-5p/hsa-miR-124-3p (Fig. 5i–n). Several other molecules involved in cancer-survival signalling, including PIK3R, RAS, MAPK, and STAT, were also significantly dysregulated upon miRNA transfection in A2780 cells (Fig. 5o–t), respectively. The validated target molecules agreed with the predicted targets associated with KEGG-specific cancer-related pathways for the miRNAs identified from CM-derived exosomes (Supplementary Fig. S8). These results suggested that overexpression of specific exosomal miRNAs dysregulated molecules related to cell-cycle progression, cytokine and cytokine-receptor expression, and cancer-survival pathways.

Internalization of hAMSC-CM-derived exosomes upregulates pro-apoptotic molecules in A2780 cells. Immunoblotting data in Fig. 6a showed upregulated expression of several pro-apoptotic signalling molecules, including phosphorylated p33, BAX, activated CASP9, and activated CASP3 in A2780 cells that had internalized CM-derived exosomes as compared with control cells. The full-length gel images are provided in Supplementary Fig. S9. In contrast, anti-apoptotic BCL2 was downregulated in these cells relative to control cells. These data indicated that internalization of CM-derived exosome resulted in activation of the intrinsic apoptosis pathway in A2780 cells.

Discussion
MSCs are important regulators of biological processes such as healing, regeneration, maintenance, and remodeling, under both normal and disease conditions. MSCs also play important roles in cancer biology; however, their ability to suppress or promote cancer remains questionable. The data presented here showed that exosomes derived from hAMSC-CM were important carriers of miRNAs involved in the regulation of certain cancer types.
We showed that treatment of A2780 cells with hAMSC-CM reduced cell viability, increased ROS production, and reduced MPP, indicating that CM derived from hAMSCs, and specifically the components released by hAMSCs, could induce deterioration of mitochondrial integrity, promote oxidative stress, and suppress cell proliferation. Moreover, treatment with CM induced S-phase arrest of the cell cycle, which is an uncommon phenomenon. In most cases, chemically induced cell-cycle arrest occurs mainly in G0/G1- and/or G2/M-phases; however, activation of the mitochondria-mediated apoptosis pathway and downregulation of CDKs could induce S-phase arrest.

Our findings revealed that overexpression of hsa-miR-124-3p, an exosomal miRNA, led to the downregulation of different CDKs, such as CDK2, CDK4, and CDK6. Additionally, other CDK family members were also predicted as targets for other exosomal miRNAs. Therefore, exosomal miRNA-mediated dysregulation of CDKs could be an explanation for S-phase arrest of the cell cycle in A2780 cancer cells. Similarily, internalization of exosomes resulted in greater decreases in proliferation, wound-repair capacity, and colony formation ability in A2780 cells compared to treatment with hAMSC-CM alone, thereby establishing that hAMSC-CM-derived exosomes are important factors related to inhibition of A2780 cell proliferation. We also observed that while treatment of A2780 cells with protease-digested exosome lysate reduced cell viability, treatment with RNase-digested exosome lysate did not suppress cell proliferation, indicating that exosomal RNAs were the sources of inhibition.

Additionally, detection of numerous known miRNAs by NGS verified their reported involvement in regulation of growth, progression, and metastasis in different cancers. Furthermore, we validated that selected miRNA target molecules were involved in regulating cell-cycle progression, cytokine and cytokine-receptor expression, and cancer survival, suggesting that hAMSC-CM-derived exosomes are carriers of cancer-regulatory miRNAs.

Exceptionally, hAMSC-CM derived exosomal miRNAs demonstrated outstanding diversity, which encompasses numerous anti-cancer miRNAs, as well as new and poorly investigated miRNAs. In contrast, miRNAs, which are frequently detected in different cancers exosomes (include hsa-miR-105, hsa-miR-214, hsa-miR-92, hsa-miR-21, hsa-miR-29, hsa-miR-9, hsa-miR-222) were either absent or showed very nominal expression in hAMSC-CM derived exosomes. Interestingly, exosomes derived from metastatic-cancers might also display high expression for anti-cancer miRNAs such as hsa-let-7, hsa-miR-23, hsa-miR-224 and hsa-miR-921, which is believed as an internal reformation technique of the cancer cells to retain malignant properties by sacking the promising threats outside. However, presence of such anti-cancer miRNAs could not resist the pro-cancer

**Figure 5.** Validation of targets for different miRNAs in KEGG 'Pathways in cancer'. (a–e) Confirmation of specific miRNA overexpression in transfected cells as compared with control cells. (f–h) Members of the CDK family found to be downregulated in hsa-miR-124-3p transfected cells. (i–n) Different cytokines and cytokine receptors targeted by different miRNAs. (o–t) Survival signaling-pathway molecules found to be downregulated following transfection of different miRNA as compared to negative-control cells. The experiments were repeated at least three times. *p < 0.05, **p < 0.01 and ***p < 0.001.
activity of metastatic-cancer-derived exosomes. Instead, metastatic-tumor-derived exosomes functions as track-
ing system for the future site of metastasis\(^65\). Specially, the cytokines receptors (such as integrin) carried by
the tumor-derived exosomes might work as antenna to recognize the potential site of metastasis, for instance,
exosomes possess integrin \(\alpha_6\beta_4\) and \(\alpha_6\beta_1\) caused metastasis of cancer to lung, while, integrin \(\alpha_v\beta_5\) resulted
metastasis to liver\(^65\). Therefore, miRNAs are not the only players behind exosomes, rather, components includ-
ing cytokines and proteins, lipids and others carried by exosomes might perform substantial role to motivate
exosomes for its terminal accomplishment.

The mechanism(s) by which exosomes derived by hAMSCs transport different molecules, including miR-
NAs, to surrounding cells/tissues is illustrated in Fig. 6b. Once internalized by cells/tissues, the newly introduced
miRNAs could target different signalling molecules related to cell cycle progression, cytokine and oncogene
expression, and cancer-related pathways, ultimately resulting cell cycle blocking, reduced viability, upregulation
of pro-apoptotic molecules, and downregulation of anti-apoptotic molecules. Considering the limited influence
on high-grade serous CAOV-3 ovarian cancer cells, which did not show any considerable change in proliferation
and wound healing capacity upon treatment with hAMSC-CM-derived exosomes, it appears that their affects
are not identical for all cancer types, also the influence might depend on the level of malignance or stage of can-
cer progression. For instance, Lin \textit{et al.}\(^66\) showed that internalization of hAMSC-derived exosomes promoted
migration and invasiveness of MCF-7 cancer cells. Therefore, our findings suggested that the regulatory role of
hAMSC-CM-derived exosomes on cancer cells is highly variable depending upon the type of cancer and sur-
rounding niche as well as the mutation and acquired-resistance status of the cancer cell lines.

In conclusion, internalized hAMSC-CM-derived exosomes inhibited the growth and proliferation of A2780
and SKOV-3 ovarian cancer cells, suggesting that the relationship between MSCs and cancer could be partially
explained by the mechanisms involved with exosome activity, given that MSC-derived exosomes contain miR-
NAs, which could be involved in cancer suppression or progression. Exosome influence could be defined by the
enriched miRNA cluster, cancer types, and the surrounding niche. While MSCs have the potential to act as both
cancer suppressors and promoters, the terminal mode of action could be determined by MSC origin, cancer type,
and cytokine expression. The phenotypic relationships observed between MSCs and various cancers might not be
rigid, and, therefore, could be determined by the cumulative effects of molecules released by MSCs. Our data pro-
vide valuable insights into these relationships; however, further extensive investigation is required to investigate
the roles of exosomal miRNAs in different cancer types.

**Materials and Methods**

**Reagents.** ExoQuick-TC exosome-isolation reagent (EXOTC50A-1), SeraMir exosome RNA-isolation
kit (RA808A-1), exosome-specific marker anti-\(\alpha\)-CD63 antibody (EXOABCD63A-1), and florescence
exosome-preparation reagent (EXDC300A-1) were from System Biosciences (SBI, Palo Alto, CA, USA).
Penicillin-streptomycin (PS) solution, trypsin-EDTA solution, RPMI medium, and \(\alpha\)-MEM media were
obtained from Life Technologies GIBCO (Grand Island, NY, USA). Fetal bovine serum (FBS) was from Sigma-
Aldrich (St. Louis, MO, USA). Antibodies used for immunoblotting were against phosphorylated p53 (Cell
Signaling Technology, Beverly, MA, USA), BAX, BCL2, CASP9, CASP3, \(\beta\)-actin (Abcam, Cambridge, MA),
and cytochrome c (ENZO Diagnostics Inc., Farmingdale, NY, USA).

**Conditional medium (CM) preparation.** hAMSCs were kindly donated by Maria Biotech Co. Ltd. (Seoul,
South Korea) and cultured in \(\alpha\)-MEM media containing 10% FBS and 100 U/mL PS in a humidified incubator at
37°C and 5% CO\(_2\). The hAMSC-CM was collected from cells passage between six and seven times. Briefly, \(2 \times 10^6\)

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**Figure 6. Potential mechanism of apoptosis activation in exosome-treated A2780 cancer cells.** (a) Exosome-
treated cells displayed upregulated expression of pro-apoptotic molecules and downregulated expression of
anti-apoptotic BCL2. (b) hAMSC-CM-derived exosome inhibition of A2780 cell proliferation. hAMSCs release
exosomes in CM, where exosomes function as carriers of MSC-secreted miRNAs. Upon internalization by
cancer cells, miRNA is released by exosomes to target genes expressing different molecules associated with
cancer proliferation, thereby disrupting cancer metastasis. The experiments were repeated at least three times.
cells were seeded per 10-cm culture dish and incubated overnight at 37 °C and 5% CO₂ for proper attachment. Media was replaced with new media, and cells were cultured for an additional 48 h, after which the resulting medium was considered hAMSC-CM. The pH stability was confirmed using both an electric pH meter and pH paper. A total of 1 L CM was collected, filtered, mixed, and aliquoted in 15-mL and 50-mL tubes for storage at −20 °C.

**CM dosage determination and cell-viability assay.** To detect the effect of hAMSC-CM on A2780 cell proliferation, a cell-viability assay was performed using 0% to 30% of CM supplementation to the fresh medium. A2780 cells were seeded (1 × 10⁴ cells/well) onto 96-well, flat-bottomed culture plates and incubated for 24 h at 37 °C in a 5% CO₂ incubator. The medium was replaced with control (fresh) or treatment (0–30% CM) medium, and cell viability was determined using a cell counting kit-8 (CCK-8; Dojindo Laboratories, Tokyo, Japan). Absorbance was measured at 450 nm in a microtiter plate reader (Multiskan FC; Thermo Fisher Scientific, Waltham, MA, USA). Based on obtained results, 25% CM supplementation to fresh medium was used for all experiments.

**ROS detection.** ROS formation was quantified using 2′,7′-dichlorodihydrofluorescein diacetate (H2-DCFDA; Sigma-Aldrich). Cells were incubated with 10 μM H2-DCFDA at 37 °C in a humidified CO₂ incubator for 30 min. Cells were then washed and re-suspended in phosphate-buffered saline (PBS), followed by detection of fluorescence emission and excitation at 515 nm and 488 nm, respectively, using a Gemini EM (SpectraMAX; Molecular Devices, Sunnyvale, CA, USA).

**MMP assessment.** MMP was detected by the cationic fluorescent indicator JC-1 (Molecular Probes, Eugene, OR, USA). Cells were incubated with 10 μM JC-1 at 37 °C for 15 min, washed with and re-suspended in PBS, and the fluorescence intensity was measured. MMP was expressed as the ratio of the fluorescence intensity of the JC1 monomers to JC1 aggregates. JC1 aggregates indicate red fluorescence for intact mitochondria, with emission at 583 nm, while JC1 monomers demonstrate green fluorescence in cytoplasm, with emission at 525 nm and excitation at 488 nm.

**Cell cycle.** A2780 cells were seeded onto 6-well plates (30,000 cells/cm²) and allowed to grow overnight, after which the medium was replaced with control (fresh) or treatment (25% CM) medium every 24 h. Cells were harvested at 48 h and 72 h for cell cycle analysis, washed with PBS, fixed with 70% ethanol, and stored at −20 °C overnight. Fixed cells were washed again with PBS, treated with RNaseA solution, and stained with propidium iodide. Cell-cycle analysis was performed using BD FACS Calibur (BD Biosciences, San Jose, CA, USA). The raw data was analysed using ModFit software trial version (http://www.vsh.com/products/mflt/index.asp).

**TMR red assay.** A2780 cells were seeded onto 6-well plates (30,000 cells/cm²) and allowed to grow overnight. Medium was replaced with control (fresh) or treatment (25% CM) medium, and the medium was replaced every 24 h. At 72 h, cells were harvested and apoptosis rate was detected using an in situ cell-death detection kit (TMR red; Roche Diagnostics, Indianapolis, IN, USA) according to manufacturer instructions.

**Immunoblotting.** Radio-immunoprecipitation buffer was used to lyse the cells, and protein concentrations were measured with the BCA protein assay kit (Thermo Scientific). Proteins were resolved using 12% SDS-PAGE and transferred electrophoretically to polyvinylidene fluoride membranes. Membranes were incubated with 5% skim milk for 2 h at room temperature (RT), then overnight at 4 °C with primary antibodies, and 1 h with horseradish peroxidase-conjugated secondary antibody at RT. Immuno-reactivity was detected by enhanced chemiluminescence reagents.

**Exosome isolation and characterization.** Exosomes were isolated according to manufacturer instructions (ExoQuick-TC kit). Briefly, ExoQuick-TC reagent was added to hAMSC-CM at a 1:5 ratio and mixed by inverting the tubes several times. The mixture incubated overnight at 4 °C, centrifuged at 1500 g for 30 min at RT, and exosomes were recovered. For TEM, an exosome-TEM grid was prepared according to manufacturer instructions (Exosome-TEM-easy kit; 101Bio, Palo Alto, CA, USA), and exosome particles were visualized using an Hitachi H7600 electron microscope (Hitachi, Tokyo, Japan) at 100 kV accelerating voltage.

**Cell-viability assay.** Exosomes derived from each mL of CM were designated as 1 U exosomes. For viability assays, cells were treated with exosomes at one U/mL. When using protease- or RNase-digested exosomes, the control medium was supplemented with equal amounts of protease or RNase solution to avoid experimental error due to the presence of protease or RNase. For TEM, an exosome-TEM grid was prepared according to manufacturer instructions (Exosome-TEM-easy kit; 101Bio, Palo Alto, CA, USA), and exosome particles were visualized using an Hitachi H7600 electron microscope (Hitachi, Tokyo, Japan) at 100 kV accelerating voltage.

**Wound-repair assay.** For detecting wound-repair capacity, cells were seeded (30,000 cells/cm²) on 12-well plates and incubated overnight at 37 °C at 5% CO₂. Wounds were made using 10-μl pipette tips, cells were washed with PBS, and fed with either control (fresh) or different treatment combinations (25% CM, 1 U/mL exosomes, or 1 U/mL exosomes plus 25% CM), and the cells were cultured for 2 weeks. Dishes containing colonies were stained with 0.05% (w/v) crystal-violet solution, including
Fluorescence-exosome preparation and internalization by cancer cells. Fluorescence exosomes were prepared and internalization of hAMSC-CM-derived exosomes by cancer cells was determined according to manufacturer protocol (EXDC300A-1, SBI, Palo Alto, CA, USA). Briefly, 50 μL of 10× Exo-Red was added to 500 μL of exosome suspension in PBS and mixed by inversion. The mixture was incubated at 37 °C for 10 min, followed by addition of 100 μL of ExoQuick-TC reagent to stop the reaction and incubation on ice for 30 min. The mixture was centrifuged for 3 min at 14,000 rpm in a microfuge, and the labelled-exosome pellet was resuspended and delivered to cancer cells. Internalization was confirmed by fluorescence microscope.

Exosomal RNA isolation. RNA was isolated from the exosome pellet using the SeraMir kit according to manufacturer protocol. Briefly, 350 μL of lysis buffer was added to 5 U exosome pellet, vortexed for 15 s, and incubated for 5 min at RT. The mixture was supplemented with 200 μL 100% ethanol, vortexed for 10 s, transferred to a spin column with collection tube, and centrifuged at 13,000 rpm for 1 min in a microfuge. The flow through was discarded, the spin column was washed twice with 400 μL wash buffer, and transferred to a new collection tube. Exosome-derived RNA was collected in 30 μL elution buffer and stored at −80 °C for downstream analysis.

RNA analysis and small-RNA library preparation. RNA concentration was measured using a NanoDrop instrument (Thermo Scientific) and sent to Macrogen (Seoul, Korea) for small-RNA sequencing. Briefly, exosomal RNA integrity was detected by Agilent Technologies 2100 Bioanalyzer (Agilent Technologies, Santa Clara, CA, USA). Small-RNA libraries were prepared by adapter ligation, reverse transcription, PCR amplification, and pooled-gel purification. Small-RNA library size was confirmed by checking the template-size distribution using a DNA 1000 chip and an Agilent 2100 Bioanalyzer (Agilent Technologies). To achieve the highest data quality, the prepared libraries were quantified using qPCR according to the Illumina qPCR Quantification Protocol Guide. We used the rapid library standard quantification solution and calculator (Roche) to generate a standard curve of fluorescence readings and calculate the library sample concentration.

Sequence-quality check and data analysis. Sequence quality was verified by FastQC (http://www.bioinformatics.babraham.ac.uk/projects/fastqc) in order to determine data quality. Unique sequences found for known miRNAs after clustering were verified using the miRBase database (www.mirbase.org/) and BLAST (http://blast.ncbi.nlm.nih.gov/Blast.cgi). Trimmed reads were considered miRNA based on having 100% identical and full-length miRNA sequences as compared with those in the miRBase database. Unmatched trimmed reads not found in miRBase were compared against the ncRNA database Rfam (http://rfam.sanger.ac.uk/). Unmatched trimmed reads to non-miRNA in the Rfam database were considered potential novel miRNAs.

miRNA detection and target prediction. Commonly expressed miRNAs among the three samples were sorted using Venny 2.0.2 BioinfoGP (http://bioinfogp.cnb.csic.es/tools/venny/). Targeted genes were predicted using the miRSystem RNA database (http://mirsystem.cgm.ntu.edu.tw/)67, which integrates seven miRNA-target gene-prediction programs, including DIANA, miRanda, miRBridge, PicTar, PITA, RNA22, and TargetScan. Target genes predicted by at least three programs were considered for further analysis.

GO and KEGG-pathway enrichment analysis. GO functional enrichment analysis was performed to verify predicted target genes against the PANTHER DNA database (http://pantherdb.org/). KEGG and BioCarta analyses were performed against the miRSystem RNA database, which integrates information from several pathway prediction programs, including KEGG (http://www.kegg.jp/)68, BioCarta (http://cgap.nci.nih.gov/Pathways/BioCarta_Pathways), Reactome (http://www.reactome.org/), and pathway interaction database (http://www.ndexbio.org/).

Validation of miRNA expression in exosomes by qRT-PCR. Sequence validation to confirm miRNA in hAMSC-CM-derived exosomes was performed by qRT-PCR according to the instructions provided with the Mir-X miRNA qRT-PCR SYBR kit (Clontech Laboratories, Inc., CA, Mountain View, CA, USA). Briefly, cDNA was prepared from exosomal RNA by single-step polyadenylation and reverse-transcription reactions, followed by quantification of miRNA expression by qPCR. Known miRNA sequences were used as miRNA-specific 5′ primers, and mRQ 3′ primers supplied with the kit were used as 3′ primers. The U6 primer was detected as control. Primer sets are enlisted in Supplementary Table S3.

Validation of miRNA targets. For validation of miRNA-target molecules in KEGG ‘Pathways in cancer’, five synthetic miRNAs were purchased based on three strategies, 1) firstly, the miRNA should have target(s) in ‘pathways in cancer’, 2) miRNA having higher ‘read count’ were prioritized 3) a representative was selected in case of families expressed multiple members. Then, selected miRNAs were separately transfected into A2780 cancer cells using lipofectamine transfection reagents. After 72 h, total RNA was isolated using a Qiagen RNeasy kit (Qiagen, Hilden, Germany). Relative mRNA expression of selected target genes was detected by qRT-PCR. Cells
transfected with 'mir Vana™ miRNA Mimic negative Control' were considered experimental controls. Primers used for RT-PCR are enlisted in Supplementary Table S3.

**Statistical analysis.** Experiments were performed in triplicate, and statistical analysis was performed using one-way analysis of variance. Significance of differences between control and treated cells was determined by Student t test. Significance was determined as p < 0.05, p < 0.01, and p < 0.001.

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Author Contributions
A.M.M.T. Reza and J.H. Kim designed the experiments and wrote the manuscript. A.M.M.T. Reza performed the experiments and H. Yasuda critically reviewed the research. A.M.M.T. Reza and Y.J. Choi analysed the data. All authors reviewed the manuscript.

Additional Information
Accession codes: High-throughput sequencing data from non-coding exosomal RNAs were deposited in the National Center for Biotechnology Information Gene Expression Omnibus database (series accession no. GSE81151).

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