Extracellular vesicles cargo from head and neck cancer cell lines disrupt dendritic cells function and match plasma microRNAs

Extracellular vesicles (EVs) are mediators of the immune system response. Encapsulated in EVs, microRNAs can be transferred between cancer and immune cells. To define the potential effects of EVs originated from squamous cell carcinoma cells on immune system response, we performed microRNA profiling of EVs released from two distinct cell lines and treated dendritic cells derived from circulating monocytes (mono-DCs) with these EVs. We confirmed the internalization of EVs by mono-DCs and the down-regulation of microRNA mRNA targets in treated mono-DCs. Differences in surface markers of dendritic cells cultivated in the presence of EVs indicated that their content disrupts the maturation process. Additionally, microRNAs known to interfere with dendritic cell function, and detected in EVs, matched microRNAs from squamous cell carcinoma patients’ plasma: miR-17-5p in oropharyngeal squamous cell carcinoma, miR-21 in oral squamous cell carcinoma, miR-16, miR-24, and miR-181a circulating in both oral and oropharyngeal squamous cell carcinoma, and miR-23b, which has not been previously described in plasma of head and neck squamous cell carcinoma, was found in plasma from patients with these cancer subtypes. This study contributes with insights on EVs in signaling between cancer and immune cells in squamous cell carcinoma of the head and neck.

As important components of the immune system, the main function of dendritic cells (DCs) is to initiate the adaptive immune response through the processing of antigens and their presentation to T cells. Immature dendritic cells (iDCs) capture and process antigens and this physiological state is characterized by high levels of chemokine receptors (CCRs) and low levels of co-stimulatory molecules\(^1\). As iDCs migrate to lymphnodes, the maturation process is initiated and they acquire antigen presenting capabilities, a physiological state characterized by the secretion of cytokines and high expression of co-stimulatory molecules such as CD80, CD86 and MHC-II (HLA-DR)\(^2\). The differentiation is essential for DCs to become antigen presenting cells and activators of the adaptive immune response\(^3\). Eventhough there are different classification systems for DCs and discussions concerning their origin, it is possible to reproduce their differentiation process in vitro using peripheral blood monocytes\(^4\)–\(^10\).

The impairment of DC function has been widely studied in cancer. In fact, antigen recognition by T cells is one of the initial steps for the immune response against cancer, and a decrease in the frequency of mature DCs (mDCs) in cancer tissue has already been described, including for head and neck squamous cell carcinomas (HNSCC)\(^11\)–\(^13\).

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HNSCC is a heterogeneous disease that comprises a variety of tumors originating from the epithelial lining the oral and nasal cavities, larynx and pharynx. In the United States new cases may reach 34,010 in 2021, with 10,850 deaths associated with cancers from the oral cavity and pharynx alone. Heavy use of tobacco and alcohol as well as human papilloma virus infection (HPV) are among the most important risk factors for this disease. Treatment is currently based on cancer staging at the time of diagnosis and includes surgery, chemotherapy and radiotherapy, with chemo and radiotherapy used either as single therapies or in association. It has been described that immunosuppression significantly contributes to the progression of this cancer type.

Among the molecular alterations described for HNSCC, microRNAs (miRNAs) have been extensively studied. MiRNAs are small non-coding RNAs implicated in gene expression regulation. They are expressed in every human tissue but may also be found in body fluids, including serum and plasma, often associated with extracellular vesicles (EVs). EVs include exosomes and microvesicles, which originate from the cell endosomal system or that are shed from the plasma membrane, respectively. In cancer, EVs and their content, including miRNAs, have been studied in the context of cancer progression and immune system modulation. For instance, EVs content transferred to DCs has been shown to alter MHC class II expression by immune cells with ensuing immune suppression. The comprehension of the complexity of cell signaling processes and immune system modulation mediated by EVs demands proper cell models and may be dependent on the studied cancer type.

In this study, monocyte-derived dendritic cells (mono-DCs) were treated with EVs derived from oropharyngeal squamous cell carcinoma (OPHSCC) and oral squamous cell carcinoma (OSCC) cell lines. We investigated if EVs derived from cell lines could serve as useful models for studying cell signaling from HNSCC cells to immune cells by comparing EVs content with plasma miRNAs from HNSCC patients. Results indicate impairment of mono-DC maturation and migration following EV internalization, and miRNAs carried by EVs were shown to match plasma miRNAs from HNSCC patients, a subset of which target biological processes associated with immune response.

Results

EVs collected from SCC cell lines culture medium are diverse in size and include miRNAs detected in plasma from HNSCC patients. The size distribution of EVs isolated from the culture media of SCC25 and FaDu cell lines was characterized by TRPS, and EVs were visualized by transmission electron microscopy. A variety of sizes, illustrating the heterogeneity of vesicles, was observed, corroborating previous reports. In addition, flow cytometry confirmed that a subset of the isolated EVs expressed the tetraspanins CD9, CD63 and CD81, the most frequently identified proteins in exosomes.

MiRNA profiling of EVs was carried out using microarray assays. We were able to detect 96 miRNAs in EVs from FaDu and 68 miRNAs in EVs from SCC25, with 31 miRNAs common to both cell lines (Fig. 2 and Supplementary Table S1). According to current literature, of the miRNAs detected within EVs, 51 from FaDu and 24 miRNAs from SCC25 had been previously detected in plasma of OPHSCC and OSCC patients, respectively, the cancer subsites from which these cell lines derive (Supplementary Table S2).

EVs are incorporated by mono-DCs. Isolation of CD14+ blood monocytes through positive selection yielded 99% purity of monocytes, as demonstrated in Fig. 3. Immature DCs (iDCs) were obtained after cultivation with GM-CSF and IL-4 for 6 days, and mature DCs (mDCs) were obtained following treatment with LPS for additional 24 h. It was possible to observe the decreasing CD14 expression and increased expression of CD209, as expected during differentiation, and increasing levels of CD80 and HLA-DR, characterizing the maturation process (Fig. 4).

We then investigated if EVs isolated from cell culture medium could be incorporated by mDCs. The detection of EVs by fluorescence microscopy in live cells is limited by the fast dynamics of fusion events and by microscopy resolution, so in order to investigate these events we used several tools from confocal microscopy. EVs were labeled with PKH67, a green fluorescent dye, and then incubated with iDCs previously labeled with CD209 (red), a dendritic cell marker (Fig. 5A). Upon addition of EVs to the cell culture, we were able to register that the PKH67-labeled vesicles were taken up by mDCs (Fig. 5B–D). In addition, sequential images using Z-stack (Fig. 5E–L) and three-dimensional images (Fig. 5M–P) corroborated that EVs were incorporated by mDCs.

EVs interfere with dendritic cells viability, adhesion and migration. We investigated whether EVs affected the viability of DCs. iDCs (day 6) and mDCs (day 7) were treated with EVs isolated from culture medium. We observed significant reduction in cell viability of mono-DCs cultivated in the presence of EVs derived from both cell lines when compared to the control (untreated) cells (Fig. 6A–D). Viable iDCs showed differences in adhesion when cultivated in the presence of EVs, with higher number of cells attached to the culture well (Fig. 6E–G), and viable mDCs cultivated in the presence of EVs adhered more firmly to the culture dish when compared to the control cells (Fig. 6H–J).

We investigated if the difference in cell adhesion detected in culture dishes could have an impact in DC migration using transwell assays. iDCs and mDCs treated with EVs showed diminished migration when compared with control (untreated) iDCs and mDCs (Fig. 6K,L). These differences were even more evident when SCC25 was placed in the lower chamber, indicating that DCs treated with EVs did not respond to the presence of the cancer cell in the lower chamber (Fig. 6K,L).

EVs impair differentiation and maturation of mono-DCs. Monocyte-derived iDCs were obtained following CD14+ cells treatment with human recombinant IL-4 and GM-CSF for six days and the maturation signal consisted on treatment of iDCs with LPS for 24 h. Both treatments were carried out either in the presence or absence of EVs derived from FaDu or SCC25 cell lines. Immunophenotyping of monocytes and mono-DCs
was based on the detection and expression levels of cell surface markers CD14, CD33, CD80, CD86, CD209 and HLA-DR by flow cytometry.

The expression levels of CD209 and CD14 were used as differentiation markers. CD14 is highly expressed in monocytes, and during the differentiation process into mono-DCs the expression of this molecule decreases. On the other hand, CD209 is not expressed in monocytes, and during the differentiation process its expression level gradually increases, being highly expressed in mono-DCs. Our results show that mono-DCs cultivated in the presence of EVs presented higher levels of CD14 when compared with untreated mono-DCs (Fig. 7A–E), while the expression of CD209 was lower in treated mono-DCs when compared with untreated cells (Fig. 7F–J), indicating alterations in the differentiation process.

The expression levels of the molecules CD80 and HLA-DR were used as markers of cell maturation. The percentage of cells expressing CD80 and the intensity of its expression as determined by MFI were lower following treatment with EVs from FaDu or SCC25 (Fig. 8A–E). Additionally, CD80 was down-regulated at the gene expression level following treatment with EVs. The percentage of cells expressing HLA-DR, on the other hand, was similar both when mono-DCs cells were cultured in the presence or absence of EVs, but the level of HLA-DR expression, as measured by fluorescence intensity, was lower when EVs were present during the maturation
Figure 2. Venn diagram showing miRNAs detected in EVs isolated from FaDu and SCC25 cell culture medium using microarray assays.

Figure 3. Flow cytometric analysis of CD14 selected monocytes used for dendritic cells differentiation. Flow cytometry dot plot analysis (A) Exclusion of doublets, gating on FSC-Area vs FSC-High; (B) sequential gate on cells expressing CD45; (C) and a final gate displaying 99% purity, almost all cells are identified as monocytes by the co-expression of CD14+ and CD33+.
signals (Fig. 8F–J). The maturation of DCs is essential for their function in the tumor microenvironment and the differences in surface markers of mono-DCs in the presence of EVs observed here suggest an effect of their content in this process, possibly contributing to the impairment of immune responses.

Gene expression alterations found in mDCs treated with EVs suggest disrupted immune responses possibly targeted by miRNAs. In order to verify which biological processes were affected following EVs uptake by DCs, gene expression levels of mDCs treated with EVs were compared with mDCs cultured in the absence of EVs. Major processes that were clearly down-regulated when cells were treated with EVs are listed in Table 1, and include lipopolysaccharide (LPS) response, inflammatory responses, chemokine and chemokine-mediated signaling and cytokine-cytokine receptor interaction. This result suggests that the antigen presenting function of these cells is impaired after the internalization of the EVs. Down-regulated genes when DCs were treated with EVs include CCL20, FASN, IL-1B, IL-6, IL-12, NFκB, NOTCH, TMEM97, TGF-β, and TNF-α (Table 1). Additionally, IL-12, TMEM97 and CCL20 were among the most down-regulated genes in DCs following treatment with EVs (Table 2). This set of genes is highlighted due to their direct role in DC function and to the fact that they are targeted by miRNAs detected within EVs in this study: miR-16, miR-23b, miR-24, miR-17-5p, miR-21, miR-152, miR-155, miR-181, and miR-212 (Table 3). Within this group of miRNAs, a subset has been previously detected in plasma from HNSCC patients, both in previous reports and in our cohort: miR-17-5p in OPHSCC, miR-21 in OSCC, and miR-16, miR-24, miR-181a circulating in plasma of both OSCC and OPSCC patients. MiR-23b, which had not been previously described in plasma of HNSCC, was found in this work in plasma of both OSCC and OPSCC patients (Supplementary Table S2).

Discussion

Disruption of DC antigen presentation machinery has been extensively studied and shown to be one of the most significant strategies used by tumor cells to guarantee their survival and cancer progression [44,55]. EVs have been directly implicated in the cross talk between cancer cells and immune cells [29,65–68].

A previous report on the effect of EVs derived from an oral SCC cell line on monocytes and DCs showed decreased expression of antigen presentation-related components and impaired cell differentiation of DCs [39]. In our study we corroborate these results for another cell line derived from oral SCC (SCC25), and add results for a cell line derived from SCC of the hypopharynx (FaDu). Additionally, using DNA microarrays, we found significant effects on biological processes associated with DC function when mono-DCs where treated with SCC25 or FaDu-derived EVs, including the diminished expression of lipopolysaccharide (LPS) response, inflammatory response, chemokine and chemokine-mediated signaling, and cytokine-cytokine receptor interaction.
Figure 5. Illustration of EV internalization by mDCs using confocal microscopy. (A) cells labeled with CD209 (red), a dendritic cell transmembrane antigen; (B) EVs stained with PKH67 (green); (C) nuclear staining with DAPI (blue); (D) Merge of CD209, DAPI and PKH67; (E–L) Sequential images using the z-stack tool of confocal microscopy depicting EVs inside the dendritic cells. Three-dimensional images of EVs and mDCs displayed in four angles of 45° displaying nuclei (blue), CD209 (red), EVs (green); (M) front view of the cells; (N) left lateral view of the cells; (O) right lateral view of the cells; (P) back view of the cells, indicating that the EVs were internalized by mono-DCs.
Our study identified miRNAs within EVs with essential roles in DC function, a subset of these miRNAs target genes found to be down-regulated in DCs cultured in the presence of EVs (Table 3). In fact, miRNAs have been shown to contribute to the inhibition of immune responses when carried by EVs, with effects on monocyte differentiation into macrophages, production of inflammatory cytokines and co-stimulatory molecules by DCs, antigen processing and presentation, MHC II molecules expression, DC maturation, migration, and increased DC death. Additionally, these miRNAs could potentially cooperate to regulate genes involved in these processes since different miRNAs carried by EVs were shown to regulate the same genes: miR-17-5p, miR-21, miR-24 and miR-181 target TNF-α, miR-17-5p and miR-24 target IL-12, and miR-155 and miR-181 regulate IL-1β.

Even though the effects observed in DCs cannot not be entirely explained by miRNAs, we showed that EVs were internalized by DCs, and found a set of miRNA gene targets down-regulated in DCs following treatment with EVs. Additionally, these gene targets were associated with specific and essential biological processes, supporting the hypothesis that the transport of miRNAs by EVs is a tumor cell signalling tool with immune modulatory objectives in HNSCC.

It is also noteworthy that we identified a set of common miRNAs between EVs derived from SCC cell lines and plasma of HNSCC patients (Supplementary Table S2), suggesting that these plasma circulating miRNAs are derived from cancer cells and are involved in cell signaling affecting distant body sites or circulating immune cells. In fact, DCs cultivated in the presence of lung cancer patient serum have shown decreased expression of CD40, CD80, CD86, MHCII, IL1 and NFκb, similar to the results we showed here.

Finally, literature reports usually address few patients and often mixed HNSCC subtypes. Here we analyzed a cohort of individuals bearing only one of two subtypes and evaluated not only disease parameters but also sample quality and limitations of the miRNA detection technique.

In conclusion, we detected miRNAs in EVs derived from HNSCC cell lines and circulating in patients’ plasma, corroborating their potential as cancer biomarkers but also their possible implication in cell signaling, and the usefulness of these cancer cell lines to study EVs and cancer cells signaling. We found differences in gene expression and surface markers of mono-DCs cultivated in the presence of EVs collected from cancer cells culture indicating that EV content, including miRNAs, affect differntiation and maturation processes, essential aspects for the proper function of dendritic cells in the tumor microenvironment.

Methods
HNSCC-derived cell lines. The cell lines SCC25, derived from a SCC of the tongue, and FaDu derived from a SCC of the hypopharynx, were used in this study. They were obtained directly from American Type Culture Collection (catalog numbers CRL-1628 and HTB-43). The cell lines were grown in a Dulbecco’s Modified Eagle’s medium/Nutrient Mixture F-12 Ham (DMEM/F12) supplemented with 10% fetal bovine serum in a humidified atmosphere of 5% CO2 and 95% air at 37 °C.

Cell culture and EVs isolation from cell supernatant. Isolation of EVs from cell culture supernatant followed the protocols described by Skog et al. and Mathivanan et al. Briefly, SCC25 and FaDu cell lines were cultivated in five independent T75 flasks until 80-90% confluence and then kept in serum-free medium for 24 h. Following this period the medium was collected (10 mL per flask, total of 50 mL per cell line), and centrifuged for 30 min at 2000×g to remove any cells or debris. This medium was then submitted to centrifugation (4 h/52,800×g) ultracentrifugation (4 h/10,000×g) (Beckman-Coulter 70Ti rotor). The supernatant was discarded and the EVs pellet was resuspended in 400 μL of PBS and stored at −80 °C. For the treatment of mono-DCs we used 10 μL or 20 μL of this EV suspension, based on the work of Wang et al. that used the volume of medium as the reference for treating cells with EVs. The analysis of the integrity and morphologic characteristics of EVs by transmission electron microscopy followed traditional protocols. Microarray analysis, EVs from cell culture supernatant were isolated using Total Exosome Isolation Reagent (from cell culture media) (Invitrogen, Thermo Fisher Scientific, Waltham, Massachusetts, USA). Briefly, 5 mL of the isolation reagent were added to the media and mixed by vortexing. The mixture was left overnight at 4 °C. On the following day, the mixture was centrifuged for 1 h at 10,000×g at 4 °C. The supernatant was discarded and pellets were resuspended in 400 μL of PBS.

EVs characterization by flow cytometry. Immunophenotyping was performed for EVs following current protocols using the following antibodies: CD9-PE (clone:209306) from R&D systems, CD63-FITC (clone:MEM-259) from USBiological and CD81-FITC (clone:JS64) from Beckman Coulter. For fluorescence background control we used the Pure Flow (Beckman Coulter), an unstained sample of EVs and a mix of fluorescent particles with different sizes (0.1–0.9 μM) Megamix-Plus FSC (Biocytex). At least 10,000 events were acquired in Moflo Astrios EQ (Beckman Coulter), using FSC-height was used for size identification, 100 μM nozzle, pressure between 15 and 20%, sample pressure 0.3 PSI higher than sheath pressure and threshold adjusted for 488 SSC. Sample acquisition was initiated only after 5 minutes after equipment stabilization. Mask was used in FSC module with P1 mask on FSC-1 and M2 mask in FSC-2. The data analysis was performed using FlowJo software (BD Biosciences, v.10.6.2).

EVs size measurement by tunable resistive pulse sensing. Size range of isolated EVs was measured, following current recommendations. With this purpose we chose tunable resistive pulse sensing (TRPS) and the qNano equipment (Izon, Cambridge, MA, USA). Nanopores NP200 and calibration beads CPC200 (at 1:1000 dilution) were used. Isolated EVs were diluted 1:20 in 0.01% Tween-20 in PBS, filtered with Filtropur S 0.2 μm (Sarstedt, Alemanija), and measured in the same conditions as the calibration beads. The measurement
miRNA profiling of EVs content. RNA was extracted from EVs using mirCURY RNA Isolation Kit—Biofluids (Exiqon, Qiagen, Vedbæk, Denmark) following protocols provided by the manufacturer. The extracted RNA was evaluated for quality and concentration using the Nanodrop One Microvolume UV–Vis Spectrophotometer (Thermo Fisher Scientific) and stored at –80 °C. For total miRNA expression profiling of EVs we used the Affymetrix GeneChip miRNA 4.0 Array (Thermo Fisher Scientific). This microarray allows for the evaluation of 2578 mature human miRNAs (miRBase version 20). Sample preparation, hybridization, staining and washing procedures followed the manufacturer's instructions FlashTag Biotin HSR RNA Labeling Kit, (Thermo Fisher Scientific). Hybridization signals were detected using the GeneChip Scanner 3000 7G (Thermo Fisher Scientific) with Affymetrix GeneChip Command Console Software parameters recommended by the manufacturer. A probe set was considered to be expressed if 50% of the samples in the dataset have DABG (Detected Above Background) values below the DABG threshold (the default DABG was set to 0.05). Array data quality analysis, data summarization and normalization (RMA + DABG) were carried out with the Transcriptome Analysis Console software v. 4.0.1 (Thermo Fisher Scientific).

Immunophenotyping of monocytes and monocyte-derived dendritic cells during differentiation and maturation processes induced in vitro. Peripheral blood was obtained from healthy volunteers. All volunteers signed a written informed consent after the study was approved by the Institutional Review Board of Albert Einstein Ethics Committee (protocol number CAEE: 86306218.2.0000.0071). Peripheral blood was diluted 1:3 with PBS and the suspended was transferred to a 15 ml conical tube containing 5 ml of Ficoll-Paque 1.077 density (GE Healthcare, Chicago, Illinois, USA) and centrifuged (30 min/500×g) without brake at 22 °C. The cells from the interface were collected, suspended in cell culture media X-vivo 15 (Lonza, Basel, Switzerland) and centrifuged again (5 min/500×g). Monocytes were separated using CD14 positive antibody attached to beads, and selected by magnetic columns (Miltenyi Biotec, Bergisch Gladbach, Germany). The selected CD14 population was dispensed into six-well plates containing X-vivo 15 medium (Lonza, Basel, Switzerland) supplemented with antibiotic-antimycotics (Gibco, Thermo Fisher Scientific). In order to generate immature monocyte-derived dendritic cells (iDCs), the cells were cultured in the presence of human recombinant IL-4 (20 ng/mL) and GM-CSF (50 ng/mL) (both from R&D Systems, Minneapolis, MN) for six days. Mature monocytes (mDCs) were obtained after iDC stimulation with LPS 100 ng/mL for 24 h (Sigma Aldrich)49. Cells were cultivated in the presence or absence of EVs. The EVs were added on day 0, the first day of monocyte culture, and during dendritic cells differentiation (iDCs), culture day 4th. The EVs were used to treat monocytes and iDCs were collected from the culture supernatant of FaDu and SCC25 cell lines and added in two different concentrations (see “Cell culture and EVs isolation from cell supernatant”) to the same medium in which monocytes and monocytes were being cultured (X-vivo 15). The EVs were filtrated using a seringe filter of 0.22 μm, before addition to the cultures.

Immunophenotyping was performed for monocytes isolation, and during mono-DCs differentiation (iDCs) and maturation (mDCs), with the following antibodies: CD33-APC (clone:PE67.6) and HLA-DR-PerCP-Cy5.5 (clone:G46.6), from BD Biosciences, CD209-FITC (clone:DCN46), CD8-APC-Cy7 (clone:SK1), CD80-PE (Clone:L307.4), CD86-PE (Clone:2331-FUN-1), CD14-PerCP-Cy5.5 (clone:M5E2). Fluorescence minus one (FMO) was used as a fluorescence background control. At least 10,000 events were acquired using FACS LSRII FORTESSA (BD Biosciences). The data analysis was performed using FlowJo software v.10.6.2 (BD Biosciences, https://www.flowjo.com). The statistical analysis was performed using the GraphPad Prism software v.7.02 (GraphPad Software, San Diego, CA, https://www.graphpad.com/scientific-software/prism).

Detection of incorporation of EVs by mature monocyte-derived dendritic cells. EVs collected from the culture supernatant of FaDu and SCC25 cell lines were diluted in two different concentrations using the same medium in which monocytes and mono-DCs were being cultured (X-vivo 15). Before using the EVs...
they were filtered using a syringe filter 0.22 µm. Cell line-derived EVs were incubated with the membrane dye PKH67 (green fluorescent cell linker, Sigma, St. Louis, Missouri, USA) according to manufacturer’s instructions. Free dye was eliminated by adding PBS to the labeled EVs, centrifuging the mixture at (1 h/150,000×g) and then carefully discarding the supernatant. The PKH67-stained EVs were incubated with mDCs for 24 h. Cells were centrifuged at (10 min/500×g), the supernatant was discarded and this procedure was repeated twice with PBS to remove the non-internalized remaining EVs. Next, cells were transferred to a glass slide by cytopsin centrifugation, and fixed with 2% formaldehyde for 1 h. Cells were further permeabilized with Triton-X100 0.05% for 30 min and incubated in blocking solution (1% BSA in PBS) and mouse anti-human CD209 (1:200, BD) primary antibody overnight at room temperature. Next, cells were washed twice and incubated with secondary

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**Figure 7.** Expression of CD14 and CD209 during dendritic cell differentiation in the presence and absence of EV. CD14 expression following treatment with FaDu cell line EVs: (A) There were no significant differences in the percentage of iDCs expressing CD14 between control (16.2% ± 1.8%) and EV treated cells (17.1 ± 5.3% and 17.9 ± 5.5%); (B) there were significant differences in the mean intensity of fluorescence (MFI) of CD14 in iDCs between the control (2884 ± 1857) and EV treated samples (5636 ± 1613 and 7605 ± 3265) depending on EV concentration. CD14 expression following treatment with SCC25 cell line EVs: (C) There were significant differences in the percentage of iDCs expressing CD14 between the control (15.5 ± 1.3%) and EV treated cells (24.0 ± 2.3% and 33.5 ± 0.3%) depending on EV concentration; (D) there were significant differences in CD14 MFI between control (2402 ± 253) and EV treated cells (3275 ± 170 and 3623 ± 111) depending on EV concentration; (E) Representative dot plots displaying iDCs expressing CD14 on day 6 of culture without treatment (CTRL) and EV-treated cells. CD209 expression following treatment with FaDu cell line EVs: (F) There were significant differences in the percentage of iDCs expressing CD209 between control (87.4 ± 3.8%) and EV treated cells (42.8 ± 10.2% and 40.5 ± 7.1%); (G) there were differences between untreated iDCs MFI (36,670 ± 802) and treated iDCs (11,543 ± 4719 and 9900 ± 2034). CD209 expression following treatment with SCC25 cell line EVs: (H) there were significant differences in iDCs expressing CD209, between the control (96.5 ± 2.1%) and treated cells (92 ± 4.8% and 87.9 ± 1.9%), depending on EV concentration; (I) there were differences between untreated iDCs MFI (62,514 ± 12,407) and treated cells (49,110 ± 5632 and 42,011 ± 1568) (n = 4); (J) Representative dot plot graphs displaying iDCs expressing CD209 on day 6 of culture without treatment (CTRL) and EV-treated cells with different concentrations of EVs. Representative results of n = 4 samples per lineage type. The statistical analysis was performed using one way ANOVA with Dunnett correction.
anti-mouse antibody coupled with Alexa-568 (1:100, Invitrogen) in blocking solution for 1 h. After final wash in blocking solution, cells were rinsed three times with PBS and mounted in 1:10 PBS diluted Fluoromount-G with DAPI (eBiosciences, San Diego, California, USA). Cells were analyzed by confocal microscopy (Zeiss 710 Confocal Microscope, Zeiss, Germany) with specific staining for CD209 (red), EVs (green), and nuclei (blue), displayed individually and overlapped. In addition, the Z-stack tool was used for sequential photos with different depth, and a 3D photo was captured in order to show the localization of EVs inside the mDC.

Cell viability analysis of mono-DCs treated with EVs. The evaluation of the viability of mono-DCs cultivated in the presence and absence of EVs was performed by trypan blue staining and cell count in Neubauer chamber. iDCs (day 6) and mDCs (day 7) cultivated in the presence or absence of EVs derived from FaDu or SCC25 cell lines were observed using phase contrast microscopy (FSX100, Olympus, Tokyo, Japan).

Migration assay. Migration of iDCs (day 6) and mDCs (day 7) cultivated in the presence or absence of EVs derived SCC25 was evaluated using a transwell assay. iDCs or mDCs were collected and centrifuged...
(5 min/500×g) and stained using CellTrace Violet BV421 (Thermo Fisher Scientific) for 15 min at 37 °C, and then washed with PBS. Following the staining, cells were seeded on the upper chamber of a transwell (5.0 µM pores) in 24 well plates. DCs migration was induced using SCC25 cells cultivation in the lower chamber of the transwell, while for controls only cell medium was used in the lower chamber of the transwell. After 24 h the percentage of cells that migrated was evaluated by flow cytometry detecting CellTrace Violet stained DCs (iDCs and mDCs) and the antibodies CD11c-PE-CF594 (clone: B-ly6) from BD—Pharmingen and CD209—PerCP-Cy5.5 (clone: DCN46) from BD—Horizon. The events were acquired using FACS LSRII FORTESSA (BD Biosciences). The data analysis was performed using FlowJo software v.10.6.2 (BD Biosciences, https://www.flowjo.com). The statistical analysis was performed using the GraphPad Prism software v. 7.02 (GraphPad Software, San Diego, CA, https://www.graphpad.com/scientific-software/prism).

Table 1. Biological processes down regulated in mDCs treated with EVs when compared with untreated mDCs. The p value was FDR corrected.

| GO term                                      | Genes                                                                 | FDR p-value |
|----------------------------------------------|-----------------------------------------------------------------------|-------------|
| GO:0006695: cholesterol biosynthetic process | TM7SF2, EBR, MSMO1, MVD, CYP51A1, HMGCGR, HMGC51, FDPS, AICL, FDF71, INSIG2, SQLE, DHCR7, INSIG1, MVK, IDH1, HSD17B7, NSDH1, DHCR24 | 1.44E-15    |
| GO:0006954: inflammatory response            | CXCL1, CIAR1, CCL2, TNF, PTGS2, ADORA2A, AIF1, FPR1, CXC128, NFKB1, NFKB2, TLRS, TLRB, CCL24, PTGDR, IL23A, REI, CCI20, CCL3L1, CCL3L3, MGLL, IL1B, ZC3H12A, PTX3, NFKBIZ, IL1A, GBPS, ADGRE2, OLR1, HCK, CD180, CCL17, RP56KAS, TNFRSF9, ORMI, TNFAIP6, P2RX7, CCL3, CCR5, CD14, CAMK1D | 2.71E-10    |
| GO:0071294 – cellular response to zinc ion   | MT1H, MT1M, MT1A, MT2A, MT1E, MT1B, MT1H, MT1X, MT1G, MT1F             | 8.96E-07    |
| GO:0071222: cellular response to lipopolysaccharide | MEF2C, IIL6, TNF, CCL2, CXCL8, NFKB1, TLRS, CD180, SRC, IL18R2, CD36, CCL20, CD80, CR5, HAMP, ZC3H12A, IL12B, CD14 | 1.63E-05    |
| GO:0008285: negative regulation of cell proliferation | CXCL1, HIST1H2AC, BARRSE1, PTGDS, ADORA2A, CXC128, MENI, CD9, PTK2B, CCL3L1, CCL3L3, IL1B, NDRG1, ETV3, DHCR24, DNA5, IL6, KLF11, PIM2, SLC39A3, SLIT3, SOD2, OSM, AT5F, NOTCH3, TNFRSF9, RASSF5, Btg2, IRF1, EFDR, TPS3, INPFP1 | 6.75E-04    |
| GO:0070098: chemokine-mediated signaling pathway | CCL24, CXCL1, CCL3, CCR5, CCL20, CMLR1, CCL3L1, PTG2, CCL12, CCL3L2, CCL17 | 3.43E-07    |
| hsa00100: Steroid biosynthesis               | TM7SF2, EBR, SC5D, MSMO1, SQLE, CYP51A1, DHCR7, HSD17B7, NSDH1, FDFT1, DHCR24 | 1.01        |
| hsa04978: Mineral absorption                 | ATP1B1, MT1M, MT1A, MT2A, MT1E, MT1B, MT1H, MT1X, MT1G, MT1F          | 0.05        |
| hsa04062: Chemokine signaling pathway        | CXCL1, CCL2, NCP1, HCK, CCL18, NFKB1, SRC, ELMO1, CCL17, CCL24, NAR5, CCL3, CCR5, CCL20, PTK2B, CCL3L1, CCL3L3, GN4B, PAK1, JAK5 | 0.04        |
| hsa04060: Cytokine-cytokine receptor interaction | CXCL1, IL6, IL1B1, TNF, CCL2, TGFBR1, TGFBR2, TNFRSF15, CCL1, CCL17, IL17RA, CCL24, OSM, TNFRSF9, CCL3L3, IL23A, CCL5, CCL26, CCL3L1, CCL3L3, CSF3R, IL1B, IL12B | 0.04        |

Table 2. Ten most differentially expressed genes between mDCs treated with EVs and untreated mDCs. The p value was FDR corrected.

| Gene symbol | Fold change: untreated DCs vs treated DCs | FDR p-value | Gene name                |
|-------------|------------------------------------------|-------------|--------------------------|
| MUCL1       | −189.9                                   | 8.42E−07    | Mucin-like 1             |
| MMP12       | −51.96                                   | 3.06E−06    | Matrix metallopeptidase 12 |
| MMP1        | −41.16                                   | 1.06E−06    | Matrix metallopeptidase 1 |
| SFINK1      | −34.75                                   | 6.96E−06    | Serine peptidase inhibitor, Kazal type 1 |
| MMP9        | −14.87                                   | 0.003       | Matrix metallopeptidase 9 |
| COL6A1      | −11.85                                   | 5.94E−06    | Collagen. type VI. alpha 1 |
| TM4SF1      | −11.4                                    | 0.0003      | Transmembrane 4 L six family member 1 |
| ABC5        | −10.69                                   | 6.96E−05    | ATP binding cassette subfamily B member 5 |
| COL6A2      | −9.93                                    | 4.86E−05    | Collagen. type VI. alpha 2 |
| TGFBI       | −7.78                                    | 1.54E−05    | Transforming growth factor beta-induced. 68 kDa |
| IL12B       | 16.06                                    | 8.63E−05    | Interleukin 12B           |
| HMGCS1      | 16.27                                    | 8.82E−06    | 3-Hydroxy-3-methylglutaryl-CoA synthase 1 (soluble) |
| CDH1        | 20.52                                    | 0.0002      | Cadherin 1. type 1        |
| HS1ST2      | 24.99                                    | 5.94E−06    | Heparan sulfate (glucosamine) 3-0-sulfotransferase 2 |
| BARRSE1     | 27.49                                    | 1.97E−05    | Retinoic acid receptor responder (tazarotene induced) 1 |
| TNFAIP6     | 30.71                                    | 8.88E−05    | Tumor necrosis factor alpha-induced protein 6 |
| IDO1        | 63.77                                    | 2.29E−06    | Indoleamine 2.3-dioxygenase 1 |
| TME4M97     | 72.65                                    | 5.94E−06    | Transmembrane protein 97  |
| ORM1        | 74.64                                    | 8.42E−07    | Orosomucoid 1             |
| CCL20       | 91.37                                    | 0.0002      | chemokine (C-C motif) ligand 20 |

(5 min/500×g) and stained using CellTrace Violet BV421 (Thermo Fisher Scientific) for 15 min at 37 °C, and then washed with PBS. Following the staining, cells were seeded on the upper chamber of a transwell (5.0 µm pores) in 24 well plates. DCs migration was induced using SCC25 cells cultivation in the lower chamber of the transwell, while for controls only cell medium was used in the lower chamber of the transwell. After 24 h the percentage of cells that migrated was evaluated by flow cytometry detecting CellTrace Violet stained DCs (iDCs and mDCs) and the antibodies CD11c-PE-CF594 (clone:B-ly6) from BD—Pharmingen and CD209—PerCP-Cy5.5 (clone:DCN46) from BD—Horizon. The events were acquired using FACS LSRII FORTESSA (BD Biosciences). The data analysis was performed using FlowJo software v.10.6.2 (BD Biosciences, https://www.flowjo.com). The statistical analysis was performed using the GraphPad Prism software v. 7.02 (GraphPad Software, San Diego, CA, https://www.graphpad.com/scientific-software/prism).
value changes in the detection of miRNAs have been reported. RNA extraction from 200 μL of plasma followed by the addition of lysis buffer was carried out according to the manufacturer’s instructions. RNA integrity was assessed using the Agilent 2100 Bioanalyzer and the RNA 6000 Nano Kit. Briefly, cells were washed with ice-cold PBS, lysis buffer was added directly to cells and the protocol was carried out according to the manufacturer’s instructions. RNA integrity was assessed using the Agilent 2100 Bioanalyzer and the RNA 6000 Nano Kit.

expression levels of miRNAs in different levels of hemolysis was performed with the aid of a hemolysis curve and literature. Detection of miRNAs in plasma of HNSCC patients. Gene expression profiling of monocyte-derived dendritic cells. We used the GeneChip Scanner 3000 7G (Thermo Fisher Scientific) with Affymetrix GeneChip Command Console Software parameters recommended by the manufacturer. Hybridization signals were detected using the GeneChip Scanner 3000 7G.

Gene expression profiling of monocyte-derived dendritic cells. RNA was extracted from mono- DC samples using the RNeasy Mini Kit (Qiagen, Hilden, Germany). Briefly, cells were washed with ice-cold PBS, lysis buffer was added directly to cells and the protocol was carried out according to the manufacturer’s instructions. RNA integrity was assessed using the Agilent 2100 Bioanalyzer and the RNA 6000 Nano Kit (Agilent Technologies, Santa Clara, CA, USA) and total RNA was stored at ~ 80 °C until use. Gene expression was analyzed using DNA microarrays Clariom S Arrays (Thermo Fisher Scientific) following the protocols provided by the manufacturer. Hybridization signals were detected using the GeneChip Scanner 3000 7G (Thermo Fisher Scientific) with Affymetrix GeneChip Command Console Software parameters recommended by the manufacturer. A probeset was considered to be expressed if 50% of the samples in the dataset had DABG (Detected Above Background) values below the DABG threshold (the default DABG was set to 0.05). Array data quality analysis, data summarization and normalization (RMA + DABG) and statistical tests for differential expression (eBayes) were carried out using the Transcriptome Analysis Console software (TAC 4.0.1) (cut-off for significance FDR corrected p-value < 0.05). Gene expression levels are presented in fold-changes. Gene Ontology (GO) term enrichment analysis was carried out using DAVID Bioinformatics Resources 6.8 and results were considered statistically significant for p-value < 0.05 after Benjamini–Hochberg False-Discovery Rate (FDR) correction.

Literature search for plasma miRNAs in HNSCC and associated with DC function. We used PubMed Central (PMC) for literature search and only articles published in English and containing the terms “Head and neck cancer”, “HNSCC” and “plasma” were selected for the correlation of miRNAs found within EVs and HNSCC patients’ plasma (Supplementary Table S2). Similarly, an association between miRNAs and processes associated with DC function found to be deregulated in this work was investigated using the search terms “dendritic cell function”, “differentiation”, “maturation”, “migration” and “dendritic cell”.

Detection of miRNAs in plasma of HNSCC patients. In order to evaluate if miRNAs found in EVs represented miRNAs circulating in plasma from HNSCC, we evaluated plasma miRNAs in a cohort of patients. Since literature reports usually address few patients and often mix HNSCC subtypes, and techniques for detection of miRNAs are very diverse, we decided to analyze a cohort of individuals bearing HNSCC from which we could evaluate disease parameters as well as sample quality and limitations of the miRNA detection technique. The cohort comprised 15 patients with confirmed anatomopathological diagnosis of oropharyngeal squamous cell carcinoma (OPHSCC) and 30 oral cavity squamous cell carcinoma (OSCC) treated at Hospital Heliopolis and Hospital das Clínicas, Sao Paulo, Brazil. This study was approved by the Hospital Heliopolis Ethics Committee, Hospital das Clínicas Ethics Committee and by the National Commission of Ethics in Research (CONEP) (protocol number CAEE: 22983313.3.0000.0071). All individuals recruited for this study provided written informed consent and the research has been performed in accordance with the Declaration of Helsinki. Tumors were staged according to the Union for International Cancer Control (UICC)/American Joint Committee on Cancer (AJCC) staging classification system for head and neck SCC (8th edition). All patients were older than 50 years at the time of sample collection, heavy smokers and had a history of chronic alcohol use. Clinical and pathological data of the individuals participating in this study are summarized in Supplementary Table S2. Peripheral blood was collected in EDTA tubes for plasma separation. The levels of free hemoglobin in the plasma samples were evaluated by spectrophotometry analysis, through the spectrophotometer NanoVue Spectrophotometer (GE Healthcare). The absorbance peaks at 414 nm are indicative of free hemoglobin. The evaluation of the expression levels of miRNAs in different levels of hemolysis was performed with the aid of a hemolysis curve that simulated possible disruptions of red cells in the plasma. For the hemolysis curve, we use different concentrations of red cells in the plasma (0.08%, 0.16%, 0.33% and 0.66% of red blood cell volume/plasma volume) (Supplementary Fig. S1). We considered values above 0.2 of absorbance as a marker of hemolysis since above this value changes in the detection of miRNAs have been reported. RNA extraction from 200 μL of plasma followed by the addition of lysis buffer was carried out according to the manufacturer’s instructions. RNA integrity was assessed using the Agilent 2100 Bioanalyzer and the RNA 6000 Nano Kit.

| miRNAs* | Target gene of miRNAs in DCs* | Reported effects on DCs* | Cell type isolated* |
|---------|--------------------------------|--------------------------|---------------------|
| miR-16-5p | FASN | Decreased differentiation | FaDu and SCC25 |
| miR-23b-5p | NOTCH and NF-κB | Decreased cytokine production | FaDu and SCC25 |
| miR-24-3p | TNF-α, IL-6 and IL-12, NF-κB | Decreased antigen processing and presentation | FaDu and SCC25 |
| miR-181a-5p | TNF-α, IL-1β, and IL-6 | Decreased cytokine production | FaDu and SCC25 |
| miR-17-5p | TNF-α and IL-12 | Decreased maturation | SCC25 |
| miR-21-5p | TNF-α, CCL20 | Decreased migration and cytokine production | SCC25 |
| miR-152-3p | TMEM97 | Decreased antigen processing and presentation | SCC25 |
| miR-155-5p | IL-1β | Increased cell death | SCC25 |
| miR-212-3p | TGF-β | Decreased antigen processing and presentation | SCC25 |

Table 3. MiRNAs detected in EVs and respective gene targets associated with immune cell function found to be down-regulated in DCs treated with EVs. Target genes reported here were studied in monocytes and/or dendritic cells according to references included in this table. *Reported in this work. *Reported in this work and literature.
the same protocol for miRNA expression profiling of EV content using the miRCURY RNA Isolation Kit—Biofluids (Exiqon). MiRNA profiling in plasma samples was carried out with Serum/Plasma Focus microRNA PCR panel (Exiqon). For cDNA synthesis we used miRCURY LNA Universal RT cDNA Synthesis Kit (Exiqon) and PCR reactions were carried out according to the miRCURY LNA Universal RT microRNA PCR System (Exiqon). The amplification was performed in 7500 Real-Time PCR System (Thermo Fisher Scientific) in 96 well plates. The amplification curves were analyzed using the 7500 software (Thermo Fisher Scientific), both for determination of Ct (Cycle Threshold) and for melting curve analysis. According to manufacturer’s instructions, we considered miRNAs as “detected” when Ct ≤ 37.

Received: 24 February 2021; Accepted: 26 August 2021

References

1. Tiberio, L. et al. Chemokine and chemotactic signals in dendritic cell migration. Cell. Mol. Immunol. 15, 346–352 (2018).
2. Théry, C. & Amigorena, S. The cell biology of antigen presentation in dendritic cells. Curr. Opin. Immunol. 13, 45–51. https://doi.org/10.1016/j.cell.2001.06.010 (2001).
3. McColl, S. R. Chemokines and dendritic cells: A crucial alliance. Immunol. Cell Biol. 80, 489–496. https://doi.org/10.1046/j.1440-1711.2002.01113.x (2002).
4. Dalod, M., Chelbi, R., Malissen, B. & Lawrence, T. Dendritic cell maturation: Functional specialization through signaling specificity and transcriptional programming. EMBO J 33, 1104–1116. https://doi.org/10.1002/embi.201488027 (2014).
5. Al-Ashmawy, G. Dendritic cell subsets, maturation and function. Dendritic Cells 2018, 11–24 (2018).
6. Allottati, A., Kotsihas, F., Magalhaes, J. G. & Amigorena, S. Dendritic cell maturation and cross-presentation: Timing matters!. Immunol. Rev. 272, 97–108. https://doi.org/10.1111/imr.12432 (2012).
7. Ardavin, C. et al. Origin and differentiation of dendritic cells. Trends Immunol. 22, 691–700. https://doi.org/10.1016/S1471-4906(01)02059-2 (2001).
8. Chapuis, F. et al. Differentiation of human dendritic cells from monocytes in vitro. Eur. J. Immunol. 27, 431–441 (1997).
9. Chometon, T. Q. et al. A protocol for rapid monocyte isolation and generation of singular human monocyte-derived dendritic cells. PLoS ONE 15, e0231132. https://doi.org/10.1371/journal.pone.0231132 (2020).
10. O’Neill, D. W. & Bhardwaj, N. Dendritic cell maturation and cross-presentation: Timing matters!. Immunol. Rev. 232, 97–108. https://doi.org/10.1111/imr.12291 (2010).
11. Gardner, A. & Ruffell, B. Dendritic cells and cancer immunity. Trends Immunol. 37, 855–865. https://doi.org/10.1016/j.it.2016.09.006 (2016).
12. Ma, Y. et al. Tumor-associated regulatory dendritic cells. Semin. Cancer Biol. 22, 298–306. https://doi.org/10.1016/j.semcancer.2012.02.010 (2012).
13. Almand, B. et al. Clinical significance of defective dendritic cell differentiation in cancer. Clin. Cancer Res. 6, 1755–1766 (2000).
14. Board, P. A. T. E. et al. Chemokine and chemotactic signals in dendritic cell migration. Curr. Opin. Immunol. 13, 45–51. https://doi.org/10.1016/j.cell.2001.06.010 (2001).
15. Board, P. A. T. E. et al. Chemokine and chemotactic signals in dendritic cell migration. Curr. Opin. Immunol. 13, 45–51. https://doi.org/10.1016/j.cell.2001.06.010 (2001).
16. Hidalgo, K. et al. Expression profiles of miR-29c, miR-200b and miR-375 in tumour and tumour-adjacent tissues of head and neck cancers. Tumour Biol. 37, 12627–12633. https://doi.org/10.1007/s12095-016-1547-2 (2016).
17. Inokihita, T., Yip, K. W., Spence, T. & Liu, F. F. MicroRNAs in extracellular vesicles: Potential cancer biomarkers. J. Hum. Genet. 62, 67–74. https://doi.org/10.1038/jhg.2016.87 (2017).
18. Valadi, H. et al. Exosome-mediated transfer of microRNAs and miRNAs is a novel mechanism of genetic exchange between cells. Nat. Cell Biol. 9, 654–659. https://doi.org/10.1038/nclb1596 (2007).
19. Fernandez-Messina, L., Gutierrez-Vazquez, C., Rivas-Garcia, E., Sanchez-Madrid, F. & de la Fuente, H. Immunomodulatory role of microRNAs transferred by extracellular vesicles. Biol. Cell 107, 61–77. https://doi.org/10.1111/boc.201400881 (2015).
20. Zaborowski, M. P., Balaj, L., Breakefield, X. O. & Lai, C. P. Extracellular vesicles: Composition, biological relevance, and methods of study. Biosciences 65, 783–797. https://doi.org/10.1010/biosci/biv6584 (2017).
21. Ding, C. et al. Pancreatic cancer-derived exosomes transfer miRNAs to dendritic cells and inhibit RFXAP expression via miR-212-3p. Oncotarget 6, 29877–29888. https://doi.org/10.18632/oncotarget.4924 (2015).
22. Corrado, C. et al. Exosomes as intercellular signaling organelles involved in health and disease: Basic science and clinical applications. Int. J. Mol. Sci. 14, 5338–5366. https://doi.org/10.3390/ijms14035338 (2013).
23. Hartjes, T. A., Mytnyk, S., Jenster, G. W., van Steijn, V. & van Royen, M. E. Extracellular vesicle quantification and characterization: Common methods and emerging approaches. Bioengineering 6, 7 (2019).
24. Bergmann, C. et al. Tumor-derived microvesicles in sera of patients with head and neck cancer and their role in tumor progression. Head Neck 31, 371–380 (2009).
25. Zhang, Z. et al. LMP1-positive extracellular vesicles promote radioresistance in nasopharyngeal carcinoma cells through P38 MAPK signaling. Cancer Med. 8, 6082–6094 (2019).
26. Fujitawa, T. et al. Anti-EGFR antibody cetuximab is secreted by oral squamous cell carcinoma and alters EGF-driven mesenchymal transition. Biochem. Biophys. Res. Commun. 503, 1267–1272 (2018).
Acknowledgements
The authors thank Dr. Roberto Cabado from Instituto de Ciencias Biomedicas, University of Sao Paulo, for technical assistance with electron microscopy and Dr. Juan Sebastian Henao Agudelo for fruitful discussions. This work was funded by FAPESP grant 10/51168-0.

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Competing interests
The authors declare no competing interests.

Additional information
Supplementary Information The online version contains supplementary material available at https://doi.org/10.1038/s41598-021-97753-y.

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