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IL-3 signalling in the tumour microenvironment shapes the immune response *via* tumour endothelial cell-derived extracellular vesicles

Tatiana Lopatina ^{a,1}, Malvina Koni ^{a,1}, Cristina Grange ^{a,1}, Massimo Cedrino ^b, Saveria Femminò ^a, Giusy Lombardo ^a, Enrica Favaro ^a, Maria Felice Brizzi ^{a,*}

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ABSTRACT

Antibody-based anti-cancer therapy is considered a successful approach to impair tumour progression. This study aimed to investigate the clinical impact of targeting the IL-3 signalling in the microenvironment of solid tumours. We intended to investigate whether the IL-3Rα blockade on tumour-derived endothelial cells (TEC) can modulate PD-L1 expression in tumour cells and peripheral blood mononuclear cells (PBMC) to reshape the anti-tumour immune response. Extracellular vesicles released by TEC after IL-3Rα blockade (aTEV) were used as the ultimate effectors of the antibody-based approach, while naive TEC-derived extracellular vesicles (nTEV) served as control. Firstly, we demonstrated that, either directly or indirectly via nTEV, IL-3 controls the expression of its receptor on TEC and PBMC respectively. Moreover, we found that nTEV, moulded by the autocrine secretion of IL-3, increased PD-L1 expression in myeloid cells both in vitro and in vivo. In addition, we found that nTEVprimed PBMC favour tumour cell growth (TEC and MDA-MB-231 cells), whereas PBMC-primed with aTEV still retain their anti-tumour properties. Isolated T-cells pre-conditioned with nTEV or aTEV and co-cultured with TEC or MDA-MB-231 cells have no effects, thereby sustaining the key role of myeloid cells in tumour immune editing. In vivo nTEV, but not aTEV, increased the expression of PD-L1 in primary tumours, lung and liver metastases. Finally, we demonstrated that the enrichment of miR-214 in aTEV impacts on PD-L1 expression in vivo. Overall, these data indicate that an approach based on IL-3Ra blockade in TEC rearranges EV cargo and may reshape the anti-tumour immune response.

1. Introduction

Tumour immune regulation leading to tumour tolerance is also orchestrated by tumour-derived extracellular vesicles (TEV) [1,2]. Extracellular vesicles (EV) are heterogenic membrane structures with a specific molecular composition dictated by their cell of origin. TEV have surface receptors and stimulating factors, such as, TLR4, HLA class I, HLA G, as well as inflammatory cytokines (TGF β 1, IFN, IL-6, IL-10) [2]. TEV are also enriched in immune relevant microRNA (miR-146, miR-214 and many others) [3–5].

TEV express tumour-associated antigens and neo-antigens conveyed by the parental tumour cells as well as major histocompatibility complex (MHC) class I and class II molecules [1]. Thus, TEV could present tumour antigens to antigen presenting cells and drive specific immune response [6]. The choice to trigger cytotoxic effect or immunosuppression

depends on the presence of co-stimulatory factors. Indeed, it has been extensively reported that TEV promote tumour immune escape through different mechanisms [1,2]. This implies that, since TEV regulate the immune response, by facilitating communication between immune cells and cancer cells, they also impact on tumour progression and metastasis formation

The programmed cell death-1 receptor (PD-1) is an immune check-point inhibitor, expressed on the surface of immune effector cells [7]. PD-1 is mainly activated by the programmed cell death ligand 1 (PD-L1), expressed by several cell types [8]. The PD-1/PD-L1 pathway fine-tunes inflammation also supporting tolerance of circulating T-lymphocyte [8]. In cancer, the expression of PD-L1 is recognized as one of the major immune escape mechanisms [7,9]. Indeed, in several cancers, PD-L1 is highly expressed and the PD-L1/PD-1 signalling is engaged to evade the T-cell-mediated immune regulation [7]. Additionally, PD-L1 was found

a Department of Medical Sciences, University of Turin, Turin, Italy

^b 2i3T Scarl University of Turin, Italy

^{*} Correspondence to: Department of Medical Sciences, University of Turin, Corso Dogliotti 14, 10126 Turin, Italy. E-mail address: mariafelice.brizzi@unito.it (M.F. Brizzi).

¹ TL, MK, and CG contributed equally

in TEV, derived from melanoma [8], prostate cancer [11], glioblastoma [10], and leukaemia [13] and has been detected in blood samples of cancer patients [14]. Recent studies have suggested that EV isolated from the blood samples of cancer patients express PD-L1, and that EV PD-L1 content correlates with patient's pathological features [12,15, 16].

Tumour endothelial cells (TEC) strictly control tumour development and metastasis formation by allowing nutrient and oxygen supply, immune cell entrance, and managing the escape of tumour cells to reach secondary sites [17,18]. TEC are unique and differ from normal endothelial cells in term of proliferation surface protein expression, secretome, and released EV [19,20]. Moreover, naive EV released by TEC (nTEV) share with TEV several activities: the ability to promote vessel formation, the capability to restrain of the immune surveillance, and the enhancement of tumour growth and metastasis formation [2,4]. We have previously shown that TEC also express the receptor for interleukin 3 (IL-3R α) [21]. More recently we provided evidence that the IL-3R α blockade on TEC changes the content of their released EV (aTEV), impairing their oncogenic action [3,4]. Due to its highly expression in leukemic cells the impact of the IL-3Rα blockade has been extensively investigated in leukaemia patients [22], while poorly explored in solid tumours. Since nTEV suppress the immune response [2,6], while the impairment of IL-3 signalling in TEC, by the receptor blockade, interferes with tumour progression, we aimed to evaluate whether and how aTEV can also revert the immunosuppressive functions of nTEV in solid tumours. Particular attention has been devoted to investigate the impact of nTEV and aTEV in the regulation of PD-L1 expression.

2. Materials and methods

2.1. nTEV and aTEV isolation

TEC were isolated as previously described [23] and grown in the complete EndoGro medium (Millipore) supplemented with 2% of fetal bovine serum (FBS). TEC were functionally evaluated to form vessel-like structures at different passages. The expression of IL-3R α on TEC untreated or stimulated with IL-3 was evaluated by Fluorescence-activated cell sorting (FACS) using anti-human IL-3R α antibody (Miltenyi Biotec, #130–113–322).

For nTEV isolation, confluent TEC were cultured in serum-free DMEM for 18 h. For aTEV collection, TEC were treated with 1 µg/ml of anti-human IL-3Rα mouse antibody (R&D Systems, #MAB301-100, Clone 32703) in serum-free DMEM. The conditioned medium from TEC untreated or treated with the blocking IL-3Rα antibody was centrifuged for 30 min at 3.000 g to remove cell debris and apoptotic bodies. Then the supernatant was filtered with PES membrane filters (0.22 µm, Millipore) and submitted to ultracentrifugation for 2 h at 100.000 g at 4 °C, using the Beckman Coulter Optima L-100 K Ultracentrifuge with the rotor type 45 Ti 45000RPM. The EV pellets were resuspended in DMEM supplemented with 1% of DMSO and stored at -80 °C until further use. After thawed, EV aliquots were resuspended in PBS and analysed using the Nanoparticle tracking analysis (NTA) by NanoSight NS300 system (Malvern Instruments, Ltd) and transmission electron microscopy (Jeol JEM 1400 Flash electron microscope, Jeol, Tokyo, Japan) [24]. For further characterisation ExoView analysis (NanoView Biosciences), and MACSPlex exosome kit (Miltenyi Biotech) were used according to manufacturer instructions.

Western blot analysis for nTEV and aTEV was performed using 7% sodium dodecyl sulfate–polyacrylamide gel electrophoresis. Immunoblotting with anti-human CD63 (Abcam, #ab134045), CD81 (Abcam, #ab109201), CD9 (Abcam, #ab223052), CD29 (Invitrogen, #Ma5–17103), and GM-130 (Abcam, #ab52649) antibody was performed overnight at 4 °C. Appropriate secondary HRP-labelled anti rabbit or mouse antibody (BioRad) were used at 1:1000 dilution for 1 h at room temperature. The protein bands were visualized with chemiluminescence (ECL) detection kit and ChemiDoc $^{\text{TM}}$ XRS + System

(BioRad). Lysates from cells, nTEV, and aTEV were loaded at concentration of $10\,\mu\text{g/well}$.

2.2. PBMC isolation and treatment with nTEV or aTEV

Fresh human PBMC were isolated by density gradient (Ficoll, Sigma) centrifugation from heparinized blood samples obtained from healthy donors. The use of PBMC was approved by the Ethic Committee of A.O. U. Città della Salute e della Scienza di Torino, Turin, Italy (CS2/1255–Protocol number 0050416, May 16, 2019). PBMC were seeded in 6-well plates at the density of $2\times 10^7/\text{well}$ in 2 ml of serum-free AIM V medium. nTEV or aTEV were added to PBMC at the concentration of $1\times 10^{10}/\text{ml}$ (approximately 1×10^3 EV/cell) for at least 24 h. As control, we used non-stimulated PBMC. After 24 h or 5 days, PBMC were counted using Muse® Count & Viability Kit (Luminex), analysed by FACS using the anti-human IL-3R α antibody (#306014, Biolegend), or used for coculture experiments or for T-cell isolation. In selected experiments, T-cells were isolated from fresh PBMC (naive) seeded at the density of $1\times 10^7/\text{well}$ in 2 ml of serum-free AIM V medium and stimulated as indicated.

For PD-L1 analysis, approximately 1×10^6 PBMC, cultured for 48 h in the presence of nTEV or aTEV were resuspended in $100\,\mu l$ of PBS supplemented with 0.1% bovine serum albumin and incubated with anti-human PD-L1 antibody (Miltenyi Biotec, #130–122–809) or PE non-immune isotypic IgG for 30 min at 4 °C. Then the cells were washed and analysed using CytoFlex from Beckman Coulter.

T-cells were isolated before or after preconditioning PBMC with nTEV or aTEV using Dynabeads $^{\rm TM}$ Untouched $^{\rm TM}$ Human kit (Invitrogen, #11344D) according to the manufacturer instructions.

2.3. ELISA assay

Relative quantification of IL-1 β , IL-10, and TGF β 1 secretion by PBMC pre-treated with nTEV or aTEV alone or co-cultured with tumour cells (MDA-MB-231 and TEC) was performed using DuoSet ELISA Development Systems (R&D Systems) according to manufacturer instructions. Naive PBMC served as internal control.

2.4. Co-culture of PBMC and TEC or MDA-MB-231 cells

To study the expression of the IL-3R α on TEC, 0.4 μm pore transwells (Costar, #3412) were used for co-culture experiments with PBMC pretreated with nTEV or aTEV. 1×10^5 TEC were seeded in 6-well plates and PBMC were stimulated with nTEV or aTEV as above described. After 24 h pre-stimulated PBMC were plated in the upper chamber of the transwell at the concentration of 5×10^6 /well and put onto TEC. After additional 48 h TEC were analysed by FACS using the anti-human IL-3R α antibody (Miltenyi Biotec, #130–113–322).

To evaluate the cytotoxic effects of PMBC against tumour, a direct cell-to-cell contact cultures were used. To this end, freshly isolated PBMC were stimulated with nTEV or aTEV as above described for 24 h, meanwhile TEC or MDA-MB-231 cells were labelled with CFSE and seeded in 6-well plates, at the concentration of $1\times10^5/\text{well}$. The day after, 5×10^6 of pre-stimulated PBMC were plated on TEC or MDA-MB-231 cells for 48 h. As control, we used TEC or MDA-MB-231 cells cultured alone or co-cultured with untreated PBMC. Six independent experiments were performed in duplicates.

2.5. Tumour in vivo models

Animal studies were conducted in accordance with the Italian National Institute of Health Guide for the Care and Use of Laboratory Animals (protocol No. 833/2020-PR). Mice were housed according to the guidelines of the Federation of European Laboratory Animal Science Association and the Ethical Committee of the University of Turin. The investigators (at least 2) were blinded when assessing the outcome. To

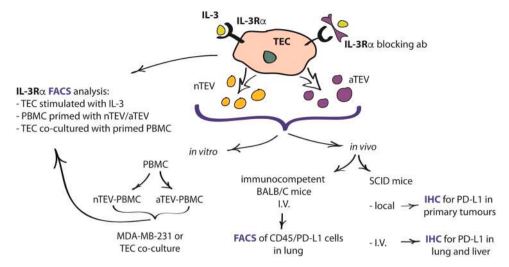


Fig. 1. Scheme of the *in vitro* and *in vivo* studies. FACS or IHC were used for PD-L1 detection. FACS analysis for IL-3Rα expression. I.V.: intravenously; IHC: immunohistochemistry.

investigate nTEV- or aTEV-mediated immune regulation *in vivo*, we used the immunocompetent female BALB/C mice. We treated mice intravenously (I.V.) with nTEV or aTEV (1×10^{10} /injection) 5 times during 2 weeks. The day after the last nTEV or aTEV injection, mice were sacrificed and PD-L1 was analysed in different tissues by FACS. To this end, the tissues were homogenized and cells positive for mouse PD-L1 (Biolegend, #124334) and CD45 (Miltenyi Biotec, #130–110–798) were detected.

Primary tumours were obtained by injecting subcutaneously Matrigel containing TEC in SCID mice (8 weeks/female) (4 mice/group) $(1\times10^6$ cells/injection). Matrigel plugs containing TEC were locally injected with saline, nTEV, or aTEV (1×10^{10}) /plug) at the third and seventh day after implantation. At day 10, the recovered plugs were embedded in paraffin (n = 4/each condition) and PD-L1 expression was analysed by immunohistochemistry, using the anti-human rabbit PD-L1 antibody (Abcam, #ab233482).

For the metastasis model, SCID mice were pre-treated intravenously with nTEV or aTEV for 5 days (1 $\times 10^{10}/\text{injection}$). On day 5, 0.6 $\times 10^6$ MDA-MB-231 cells were injected I.V. The mice were sacrificed after 5 weeks and lung and liver tissues analysed for PD-L1 expression. Liver metastases were also counted according to human PD-L1 expression.

2.6. Immunohistochemistry for PD-L1 detection in the lung and liver

Immunohistochemistry was performed using an automated slide-processing platform (Ventana BenchMark AutoStainer, Ventana Medical Systems), with Universal DAB Detection Kit detection systems. The anti-human PD-L1 rabbit polyclonal antibody (Abcam #ab233482) was diluted 1:100. Secondary HRP-labelled anti-rabbit antibody (Goat Anti-Rabbit IgG (H+L)-HRP Conjugate, BioRad) was used at 1:1000 dilution for 1 h at room temperature. Sections were counterstained with haematoxylin, dehydrated, and mounted. Quantification of the PD-L1 expression was performed using Fiji software (ImageJ). The results were expressed as mean \pm SD of PD-L1 positive area (related units) per sample (10 images/section, 4 samples/each condition).

2.7. miR-214 enrichment in nTEV

To obtaining nTEV enriched in miR-214, TEC growing in 75 cm cultured flasks were transfected with HiPerfect reagent (Qiagen) using $20\,\mu\text{M}$ hsa-miR-214 Pre-miR^M miRNA Precursor (PM12124, Invitrogen) according to manufacturer instructions. TEC transfected with scramble mimic were used as control. The day after transfection fresh FBS-free DMEM was replaced for additional 24h and collected for nTEV_miR-

214 or Scramble miRNA_TEV isolation. We used PCR to confirm the enrichment of miR-214 in nTEV_miR-214. Briefly, single-stranded cDNA was generated from total RNA sample (80 ng) by reverse transcription using miScript Reverse Transcription Kit (Qiagen) following the manufacturer's protocol. miR-214 content was measured by qRT-PCR using the miScript SYBR Green PCR Kit (all from Qiagen). All reactions were performed using an Applied Biosystems 7900HT real-time PCR instrument and run in triplicate (3 ng of cDNA for each reaction) as described by the manufacturer's protocol (Qiagen). Relative miR-214 expression was normalized to the mean expression value of RNU6 and actin housekeeping genes. Data were analysed using Expression Suite Software (ThermoFisher).

2.8. Statistical Analysis

All data are reported as mean \pm SD. Comparison between two groups was analysed by Student's t-test. Data passed both normality and equal-variance tests. One-way ANOVA followed by Tukey's multiple-comparison test was used for comparison among 3 or more groups; p-value < 0.05 was considered as significant. All $in\ vitro$ or $in\ vivo$ data are representative of at least 4 independent experiments. Graph Pad Prism version 5.04 (Graph Pad Software) was used for all statistical analyses.

3. Results

3.1. nTEV and aTEV characterisation

Firstly, nTEV and aTEV were characterized (see the scheme of the study, Fig. 1). As shown by TEM and NanoSight (data not shown), nTEV and aTEV did not differ in size (Supplementary Fig. S1). ExoView (NanoView Biosciences), MACSPlex (Miltenyi Biotech), and Western blot analyses, revealed a similar pattern of surface marker expression (CD9, CD29, CD63, and CD81) (Supplementary Fig. S1). We failed to detect PD-L1 on both nTEV and aTEV (data not shown).

3.2. IL-3R α expression in TEC

This study has been designed to investigate the potential "clinical" impact of targeting TEC using a blocking IL-3 receptor antibody, and, in particular, the effect on PD-L1 expression. Therefore, the expression of IL-3R α was first evaluated on naive or IL-3 stimulated TEC. Moreover, since IL-3 is mostly released by activated T-cells [24], to mimic the TME, IL-3R α was also analysed on TEC stimulated with nTEV-primed PBMC (Fig. 1). As shown by FACS analysis, IL-3 significantly up-regulated the

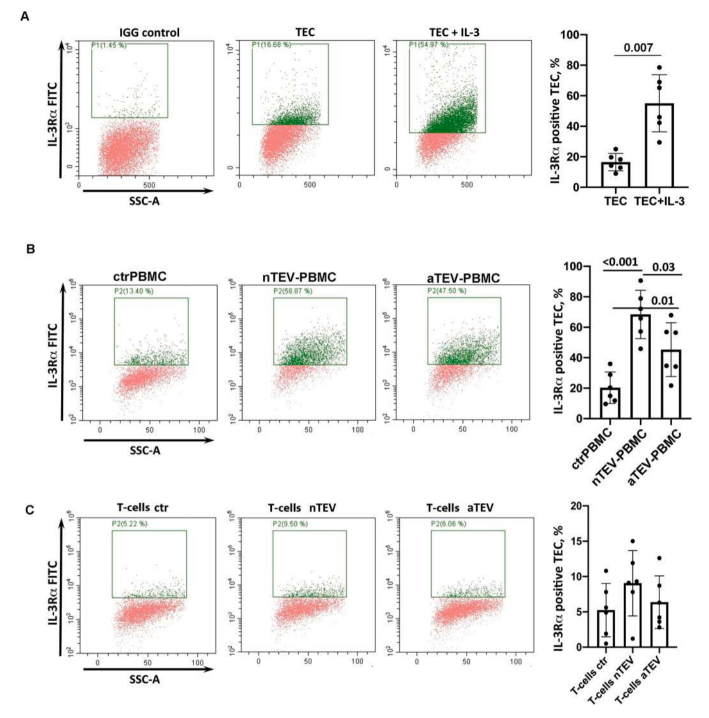


Fig. 2. IL-3R α is up-regulated in TEC co-cultured with primed PBMC and T-cells, and stimulated with IL-3. (A) Representative FACS dot plots of untreated or IL-3 stimulated TEC (24 h). (B) Representative FACS dot plots of TEC co-cultured with control PBMC (ctrPBMC) or PBMC primed with nTEV or aTEV (24 h). (C) Representative FACS dot plots of TEC co-cultured with unstimulated T-cells primed with nTEV or aTEV (24 h). Data are presented as the mean \pm SD, n=6.

expression of its receptor on TEC (Fig. 2A). No differences in PBMC viability at both 24 h and 5 days, as well as in IL-3R α expression (24 h) upon nTEV and aTEV priming were detected (Supplementary Fig. S2A-B).

Then we evaluated the expression of IL-3R α in TEC co-cultured with nTEV or aTEV-primed PBMC. We demonstrated that PBMC primed with nTEV, but not with aTEV, significantly increased the expression of the IL-3R α (Fig. 2B). Interestingly, naive T-cells (isolated from naive PBMC) pre-treated with nTEV or aTEV failed to induce the expression of IL-3R α on TEC (Fig. 2C), suggesting a key role of myeloid cells in the transfer of nTEV signals.

3.3. PBMC primed with aTEV retain their anti-tumour properties

TME immunosuppression also stems from EV-mediated communications [1]. Therefore, the effect of nTEV and aTEV in shaping PBMC anti-tumour action was further investigated. To this end, PBMC were pre-stimulated with nTEV or aTEV for 24 h, and the secretion of inflammatory factors was first analysed. We found that nTEV significantly enhance the secretion of IL-10 and TGF β 1, while aTEV increase the release of IL-1 β (Fig. 3 A). We then investigated the impact of PBMC, either untreated or primed with nTEV or aTEV, in co-cultures (direct contact) with CSFE-labelled MDA-MB-231 cells or TEC for 48 h. As

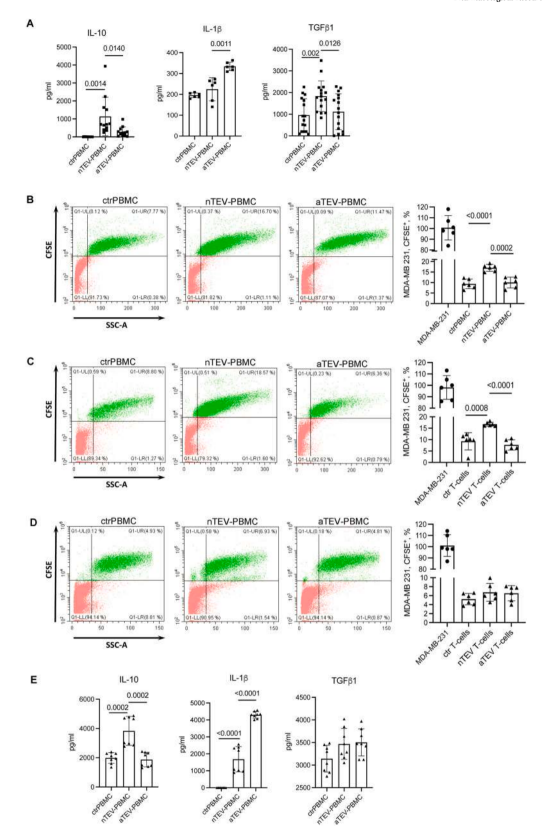


Fig. 3. nTEV, unlike aTEV, primed PBMC are impaired in their cytoxic activity. (A) Diagrams of IL-1 β , IL-10, and TGF β 1 secretion by naive (ctrPBMC) or PBMC pretreated with nTEV or aTEV. (B) Representative FACS dot plots of MDA-MB-231 cells co-cultured with ctrPBMC or PBMC pre-treated with nTEV or aTEV. Diagram represents data on MDA-MB-231 cells cultured alone (circles) or with the aforementioned PBMC (triangles). (C) Representative FACS dot plots and diagram of MDA-MB-231 cells co-cultured with T-cells, isolated from ctrPBMC or from PBMC pre-treated with nTEV or aTEV. (D) Representative FACS dot plots and diagram of MDA-MB-231 cells co-cultured with naive T-cells (isolated from ctrPBMC) pre-treated with nTEV or aTEV. (E) Diagrams of IL-1 β , IL-10, and TGF β 1 secretion by ctrPBMC or PBMC pre-treated with nTEV or aTEV co-cultured with MDA-MB-231 cells. Diagram data are presented as the mean \pm SD (at least n = 6 independent experiments).

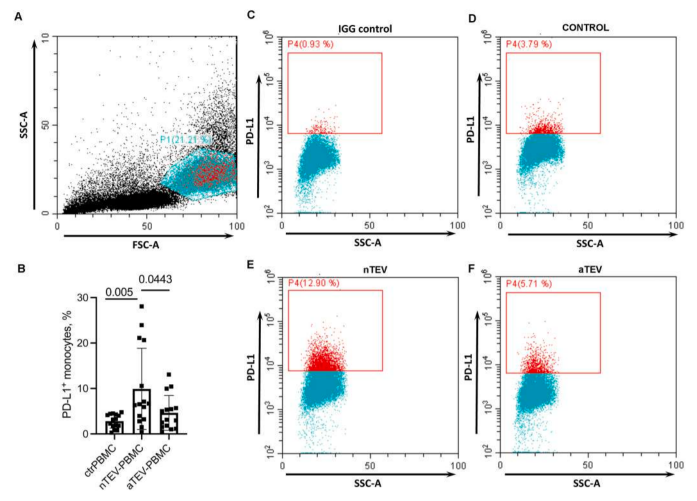


Fig. 4. nTEV, unlike aTEV, upregulated PD-L1 expression on PBMC. (A) Representative FACS dot plot image showing PBMC gating. (B) Diagram of PD-L1 expression on PBMC. Data are presented as the mean \pm SD, n=15 independent experiments. (C-F) Representative FACS dot plot images of ctrPBMC incubated with IGG control antibodies (C), ctrPBMC (D), nTEV primed PBMC (E), and aTEV primed PBMC (F) stained with anti-human PD-L1 PE conjugated antibody. In red are indicated PD-L1+ cells.

shown in Fig. 3B and Supplementary Fig. S3A we demonstrated that, while the percentage of living cells decreases in the presence of PBMC, the cytotoxic activity of nTEV primed PBMC was reduced compared to both naive and aTEV primed PBMC. To investigate whether changes in PBMC secretome may contribute to the different immune response, ELISA assay was also performed on these conditioned media. The results reported in Fig. 3D and Supplementary Fig. S3E revealed that in the presence of nTEV primed PBMC IL-10 secretion was increased. Conversely, and consistent with functional data, a reduced secretion of IL-10 and an increased level of IL-1 β were detected in co-cultures containing aTEV primed PBMC. Of note, the release of TGF\$\beta\$1 did not change (Fig. 3E, Supplementary Fig. S3E). We also investigated the contribute of T-cells. The role of T-cells in mediating the cytotoxic effect was confirmed by the observation that T-cells isolated from aTEV-primed PBMC were still able to exert anti-tumour effect when seeded together with MDA-MB-231 cells (Fig. 3 C) and TEC (Supplementary Fig. S3B). Conversely, no effect was detected when naive T-cells primed with nTEV or aTEV were co-cultured with MDA-MB-231 cells and TEC (Fig. 3D, Supplementary Fig. S3C). These data indicate that nTEV contribute to suppress the immune response and myeloid cells are key supervisors.

3.4. aTEV fine-tune PD-L1 expression

To evaluate if the expression of PD-L1 in myeloid cells may

contribute to our observations, FACS analysis was first performed on PBMC primed with nTEV or aTEV. We found that, unlike aTEV, nTEV significantly increased the expression of PD-L1 on myeloid cells (Fig. 4).

Since no data are so far available on the role played by nTEV and, most importantly, by aTEV in an immunocompetent context, *in vivo* experiments were performed in BALB/C mice intravenously injected with nTEV or aTEV for 2 weeks (Fig. 5 A). PD-L1 and CD45 coexpression were evaluated in lung, bone marrow, spleen, and peripheral blood by FACS analysis. As shown in Fig. 5B, the number of PD-L1 $^+$ /CD45 $^+$ cells was significantly increased in the lung of nTEV-treated animals, but not in spleen, bone marrow, and peripheral blood (data not shown). Of note, we found that aTEV *in vivo* administration reduced the percentage of PD-L1 $^+$ /CD45 $^+$ cells isolated from the lung.

Since tumour cells also express PD-L1 and contribute to immune evasion, the ability of nTEV and aTEV to regulate its expression on TEC derived tumours was further evaluated. We demonstrated that intratumour injection of nTEV in Matrigel plugs containing TEC enhanced PD-L1 expression, while aTEV significantly reduced its expression (Fig. 5 C,D). This indicates that nTEV and aTEV can control PDL-1 expression in different cell population in the TME, including, at least, myeloid cells and tumour cells.

Finally, to investigate the possibility that aTEV may also impact on tumour cell recruitment by regulating PD-L1 expression on homed cells, nTEV and aTEV were injected intravenously for 5 consecutive days before MDA-MB-231 cell administration (Fig. 6 A). The expression of

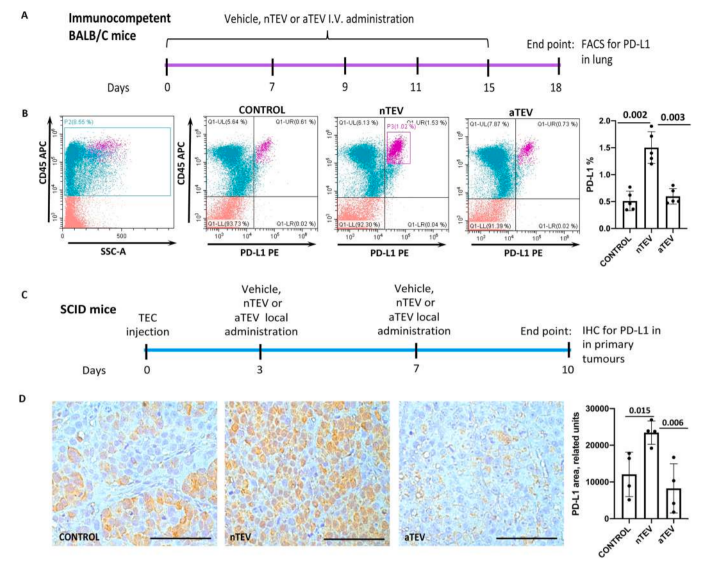


Fig. 5. nTEV, but not aTEV, increase the expression of PD-L1 in the lung and primary tumours. (A) Schematic representation of the *in vivo* study. (B) Representative FACS dot plot images showing CD45 $^+$ /PD-L1 $^+$ lung-derived cells from control, nTEV, and aTEV treated mice. Data are presented as the mean \pm SD, n = 5. (C) Schematic representation of the *in vivo* study in SCID mice. (D) Representative images of Matrigel containing TEC treated with nTEV, or aTEV stained with PD-L1 antibody. Saline treatment served as control. Scale bars = 100 μ m (original magnification, 400X). Diagram data are presented as the mean \pm SD, n = 4. I.V.: intravenously; IHC: immunohistochemistry.

PD-L1 was analysed in the lung and liver of immune-deficient animals. As shown in Fig. 6B-C, PD-L1⁺ area were significantly increased in mice primed with nTEV both in the lung and liver tissues. Again, in aTEV treated animals a significant reduction of PD-L1⁺ area was detected in both organs. Finally, as previously reported in the lung [4] a reduced number of metastatic foci was detected in the liver of aTEV primed animals (Fig. 6 C).

This observation provides evidence that aTEV priming can prevent the recruitment/homing of PD-L1 $^{+}$ tumour cells both in the lung and in the liver. More importantly, these results indicate that disturbing the IL-3 signalling on TEC drives changes on their released aTEV, impacting on the expression of PD-L1 in both tumour and CD45 $^{+}$ cells.

3.5. miR-214 enrichment in aTEV regulates the expression of PD-L1

We have previously shown that aTEV miRNA content is relevant for their biological action [3,4]. It has been shown that miR-214 post-transcriptionally regulate PD-L1 expression in B-cell lymphoma [25]. Since miR-214 was found enriched in aTEV [3], we south to determine whether miR-214 enrichment could also control PD-L1 expression in our

model. To this end, the effect of nTEV enriched in miR-214 (nTEV_miR-214) was evaluated *in vivo*. Matrigel plugs containing TEC were therefore locally injected with nTEV, aTEV, or nTEV_miR-214. Similarly to aTEV, nTEV_miR-214 treatment reduced the number of PD-L1⁺ cells in TEC formed tumours (Fig. 7) compared to nTEV. These data further confirm the role of miR-214 in the regulation of PD-L1 expression [25].

4. Discussion

In the present study, we have shown that IL-3 signalling in TEC regulates the release of pro-metastatic nTEV that suppress the immune response by enhancing PD-L1 expression on tumour and myeloid cells. Conversely, an approach based on IL-3R α blockade on TEC led to the release of aTEV enriched in miR-214 that retune the aberrant antitumour immune response reshuffling PD-L1 expression. Overall, these findings provide evidence that blocking IL-3 signalling in the TME amends PD-L1 expression and reshapes the anti-tumour immune response.

Blocking the interaction between PD-1 and PD-L1 became a revolution in cancer therapeutic approaches, particularly for lung, kidney, and

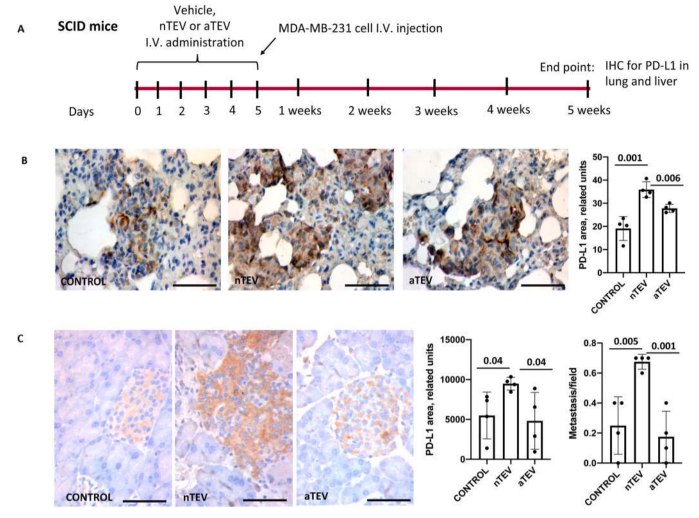


Fig. 6. PD-L1 expression is increased in the lung and liver tissues of SCID mice systemically primed with nTEV. (A) Schematic representation of the *in vivo* study. (B) Representative images and diagram of PD-L1 expression in lung tissues of control mice, and mice primed with nTEV or aTEV. Scale bars = $100 \, \mu m$ (original magnification, 400X). Diagram data are presented as the mean \pm SD, n = 4. (C) Representative images of PD-L1 expression in liver tissues of control mice, and mice primed with nTEV or aTEV. Scale bars = $100 \, \mu m$ (original magnification, 400X). Diagram data are presented as the mean \pm SD, n = 4.

bladder cancer, melanoma, and even for breast cancer [26–29]. However, the rate of success of PD-1/PD-L1 blockade in solid tumours is relatively low and mostly accompanied by both immune-related side effects [30] and TME-mediated acquired resistance [31,32]. Herein, an additional mechanism regulating PD-L1 expression in the TME is reported. In particular, we demonstrated that IL-3 signalling in TEC translates in the release of EV displaying immunosuppressive properties.

IL-3 is a hematopoietic factor mostly produced by activated T-cells, but also by monocytes/macrophages and other cell types [24]. According to proteinaltas.org (https://www.proteinatlas.org/ENS-G00000185291-IL3RA/tissue) the IL-3R α is detected in cerebral cortex and fallopian tubes in healthy subjects, while in several tumour types in cancer patients (https://www.proteinatlas.org/ENSG00000185291-IL3 RA/pathology), thus suggesting the relevance of the IL-3 signalling in solid cancer. IL-3, and in particular its binding subunit, the IL-3Rα, is a well-established acute leukaemia therapeutic target, while its role in solid tumours is yet largely undetermined [33]. The observation that the IL-3Rα is highly expressed in leukaemia stem cells, compared to their normal counterpart, has provided the rational to specifically target tumour cells [22] using an anti-IL-3Rα blocking antibody [34]. In the present study, we demonstrated that a microenvironment containing IL-3 generates a positive loop involving TEC and PBMC resulting in the up regulation of the IL-3R α in both cell types. This suggests that, as in the leukemic microenvironment, in the TME of solid tumours, the

up-regulation of IL-3R α may represent a valuable therapeutic target. Therefore, since TEC may be considered the gate for antibody entry in the TME, an antibody-based approach was used to target IL-3R α on TEC. By means of this approach we investigated whether blocking the IL-3 signalling may "educate" TEC to rearrange the cargo of their derivatives, aTEV, thereby refining the anti-tumour immune response.

TEV are involved in key aspects of cancer growth, metastatic spread, tumour immune editing, thereby considered as relevant anti-tumour targets [2,4,35]. The regulation of TEV molecular composition is highly sensitive and specific. Protein, RNA, and lipid content changes in response to microenvironment cues. We have previously shown that blocking the IL-3 signalling in TEC led to a complex rearrangement of EV molecular composition. This results in the inhibition of their pro-oncogenic functions, according to their angiogenic and metastatic actions [3,4]. Consistent with our previous results, using TEV from tumours of different origin [2], we demonstrated that nTEV impair the anti-tumour effect of PBMC against both TEC and MDA-MB-231 cells. Notably, we demonstrated that the anti-tumour effect of PBMC was retained by aTEV-primed PBMC, indicating that IL-3Rα blockade may also act on PBMC to re-establish their cytotoxic activity. Moreover, unlike T-cells isolated after PBMC pre-conditioning naive T-cells, directly stimulated with nTEV or aTEV, have no effect on tumour cells. This observation strongly supports the notion that nTEV-mediated immune regulation mainly involves mechanisms orchestrated by

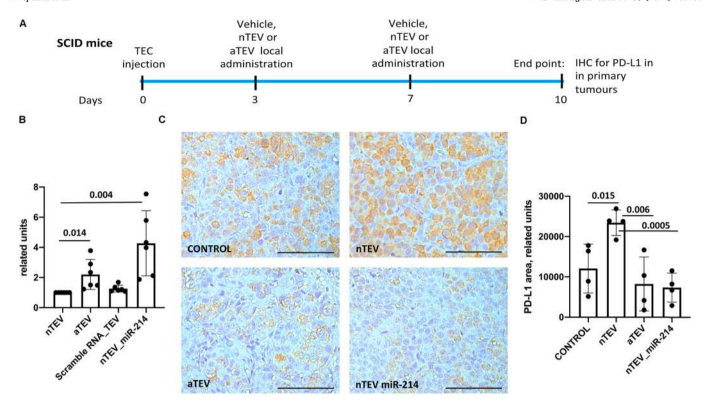


Fig. 7. miR-214 controls PD-L1 expression in vivo. (A) Schematic representation of the in vivo study. (B) Diagram of PCR analysis of miR-214 expression in nTEV, aTEV and nTEV_miR-214. As a control Scramble RNA was used. Data are presented as the mean \pm SD, n=6. Representative images (C) and diagram (D) of PD-L1 staining on TEC in control tumours, and in tumours from mice locally treated with nTEV, aTEV or nTEV_miR-214. Scale bars = 100 μ m (original magnification, 400X). Diagram data are presented as the mean \pm SD, n=4.

antigen-presenting cells. As a matter of fact, we demonstrated that nTEV, but not aTEV, increased the expression of PD-L1 on primary myeloid cells *in vitro* and *in vivo* in an immunocompetent model. This observation provides the first evidence that nTEV can control PD-L1 expression in an immunocompetent context.

PD-L1 is expressed by different tumour cell types [36], and its overexpression represents one of the most relevant mechanisms of T-cell exhaustion [37] and tumour progression. We herein demonstrated that intra-tumour nTEV injections was able to increase the expression of PD-L1 on tumour cells, and, more importantly, that aTEV bring back PD-L1 expression to the control level. Using a different mouse model, we also demonstrated that nTEV and aTEV can differently modulate PD-L1 expression in tumour cells engaged into metastatic sites. Of note, in this model we found that the expression of PD-L1 correlated with the number of metastases both in the lung and in the liver. Therefore, these results demonstrate that an antibody-based approach against the IL-3R α may reshape PD-L1 expression and the anti-tumour immune response, via aTEV.

Different mechanisms control PD-L1 expression in cancer [38]. Inflammatory cytokines released in the TME have been reported to boost the expression of PD-L1 in tumour and stromal cells [39,40]. We herein demonstrated that the inflammatory cytokine IL-3, released in the TME, besides controlling the tumour vasculature takes part in cancer immune escape, *via* nTEV. Moreover, the observation that IL-10 was increased when PBMC were primed with nTEV and left in co-cultures with cancer cells, sustains the role of nTEV in the reshuffle of the TME secretome and in tumour immune suppression. The ability of aTEV to reverse PBMC secretome and to re-establish PBMC-mediated anti-tumour cytotoxic activity, further supports the role of IL-3 in tumour immune tolerance.

Several studies have designated EV as PD-L1 delivery system [10]. However, we failed to detect PD-L1 in nTEV suggesting that, rather than transferring PD-L1 to immune or tumour cells, nTEV tune the adaptive immune response by a mechanism involving their cargo.

Epigenetic mechanisms such as histone acetylation/methylation or abnormal miRNA expression are instrumental for cancer immune escape *via* PD-L1 expression [25,38]. miR-200, miR-326, miR-34a and miR-214 are from among the most relevant miRNAs involved in PD-L1 expression [25,38,42]. Consistent with these data, we demonstrated that the enrichment of miR-214 in aTEV down-regulates PD-L1 expression and may conceivably break down tumour immune evasion. However, since EV action rely on their entire cargo we cannot rule out the possibility that cancer microenvironment can hijack the TEV cargo at different level to facilitate discrete mechanisms of the multi-step cancer process.

5. Conclusions

Overall, we provide evidence that interfering with the IL-3 signalling, by blocking the activity of its receptor at the entrance of the TME, may be instrumental for reschedule the anti-tumour immune response. Downregulating PD-L1 expression in myeloid cells also improves the response of combined immune checkpoint approaches [43]. Therefore, our findings offer a new therapeutic window to boost and reshape the aberrant immune response in solid tumour.

CRediT authorship contribution statement

TL was involved in conceptualization, methodology, data curation and writing the original draft; MK provided formal analysis of *in vitro* and *in vivo* experiments; CG performed *in vitro* and *in vivo* experiments and provided formal analysis of the *in vitro* and *in vivo* studies; MC performed the animal studies; SF isolated and characterized extracellular vesicles; GL performed the animal study; EF performed *in vitro* studies; MFB was involved in conceptualization, supervision, writing, review and editing the Ms, as well as in providing resources.

Declaration of Competing Interest

The authors declare no conflict of interest.

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Appendix A. Supporting information

Supplementary data associated with this article can be found in the online version at doi:10.1016/j.phrs.2022.106206.

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