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RAPID COMMUNICATION

Rapid communication: insights into the role of extracellular vesicles during Auger radioimmunotherapy

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\textbf{ABSTRACT}

\textbf{Purpose:} Non-targeted effects, including bystander and systemic effects, play a crucial role during Auger targeted radionuclide therapy. Here, we investigated whether small extracellular vesicles (sEVs) produced by irradiated cells could contribute to the bystander cytotoxic effects in vitro and also to therapeutic efficacy in vivo, after their injection in tumor xenografts.

\textbf{Materials and methods:} B16F10 melanoma donor cells were exposed to radiolabeled antibodies (Auger radioimmunotherapy, RIT) for 48 h or to X-rays (donor cells). Then, donor cells were incubated with fresh medium for 2 h to prepare conditioned medium (CM) that was transferred onto recipient cells for bystander effect assessment, or used for sEVs enrichment. Resulting sEVs were incubated in vitro with recipient cells for determining bystander cytotoxicity, or injected in B16F10 melanoma tumors harbored by athymic and C57BL/6 mice.

\textbf{Results:} In vitro analysis of bystander cytotoxic effects showed that CM killed about 30–40\% of melanoma cells. sEVs isolated from CM contributed to this effect. Moreover, the double-stranded DNA (dsDNA) content was increased in sEVs isolated from CM of exposed cells compared to control (not exposed), but the difference was significant only for the X-ray condition. These results were supported by immunodetection of cytosolic dsDNA in donor cells, a phenomenon that should precede dsDNA enrichment in sEVs. However, sEVs cytotoxicity could not be detected in vivo. Indeed, in athymic and in immunocompetent mice that received four intratumoral injections of sEVs (1/day), tumor growth was not delayed compared with untreated controls. Tumor growth was slightly (not significantly) delayed in immunocompetent mice treated with sEVs from X-ray-exposed cells, and significantly with sEVs purified from CM collected after 48 h of incubation. These results highlight the need to determine the optimal conditions, including radiation absorbed dose and sEVs collection time, to obtain the strongest cytotoxic effects.

\textbf{Conclusions:} This study demonstrates that sEVs could play a role during Auger RIT through bystander effects in vitro. No systemic effects were observed in vivo, under our experimental conditions. However, X-rays experiments showed that sEVs collection time might be influencing the nature of sEVs, a parameter that should also be investigated during Auger RIT.

\textbf{Introduction}

Atom decay via internal conversion or electronic capture creates vacancies in the orbital inner shells that can be filled by transitions of electrons from the outer shells. These transitions are accompanied by emission of X-rays or low-energy Auger electrons, Coster–Kronig and super Coster–Kronig electrons, depending on the involved electron transitions; however, all the ejected electrons are usually called Auger electrons (AEs). Most AEs have energy below 1 keV with very short range in tissue (<1 \textmu m) (Pomplun et al. 1987; Kassis 2004; Howell 2008) and high linear energy transfer (4–24 keV/\textmu m), compared with gamma rays and beta particles (0.2 keV/\textmu m). As nuclear DNA is the most sensitive target in cells, many studies investigated the best way to drive AEs into cell DNA (Rosenkranz et al. 2020).

This was satisfyingly obtained with drugs that behave like chemotherapy, for example radioactive DNA base analogs (e.g. 5-\textsuperscript{[125]}I抉择-2’-deoxyuridine; \textsuperscript{[125]}I-UdR) (Kassis et al. 1987). Yet, the development of more targeted approaches in cancer therapy requires to target specifically cancer cells. This was obtained using peptides and monoclonal antibodies (mAbs) (Costantini et al. 2007; Pouget et al. 2008; Santoro et al. 2009; Aghelvia et al. 2017; Ku et al. 2019). However, these molecules generally recognize cancer cell surface receptors and are internalized in endosomes where they are degraded or recycled before they can reach the nucleus (Rosenkranz et al. 2020). About 10 years ago, we demonstrated in vitro and in mice that AEs display some antitumor efficacy when targeted to the membrane of tumor cells using mAbs. We showed that targeting iodine 125...
(\(^{125}\)I) to nuclear DNA using \(^{125}\)I-UdR was a more efficient approach and killed about 50% of cells compared with 15% when \(^{125}\)I-mAbs were targeted to the cell membrane. However, we also found that about 30% of cells were killed by bystander effects (Paillas et al. 2013; Paillas et al. 2016). Therefore, we concluded that bystander effects could compensate for the anticipated inferior efficacy of the absence of nuclear targeting. Moreover, AEs-labeled mAbs bypass the disadvantages of radioactive chemotherapy. We demonstrated that cell membrane-mediated targeted and non-targeted (bystander) effects were associated with lipid raft formation and subsequent activation of signaling pathways involved in cell death (Ladjohounlou et al. 2019). However, we did not investigate the nature of the factors involved in \(^{125}\)I-mAb-induced bystander effects.

Extracellular vesicles (EVs) are one of the most attractive and promising candidates identified in the last decade for participating to intercellular communications. EVs are commonly classified in three subcategories: (i) apoptotic bodies (diameter range 1–5 \(\mu\)m) produced during apoptosis; (ii) microvesicles (100 nm–1 \(\mu\)m) (MVs), which bud directly from the plasma membrane; (iii) exosomes, the smallest EVs (50–150 nm), which are released by the fusion of multivesicular endosomal bodies with the plasma membrane (van Niel et al. 2018; Malloci et al. 2019). Many studies using powerful analytical tools have recently highlighted the complexity and heterogeneity of these vesicles, and revealed the existence of subtypes, depending on several parameters (cell source, state, isolation protocols, etc.) (Colombo et al. 2014; Kowal et al. 2016; Théry et al. 2018; Malloci et al. 2019). In this context, we adopted international EVs community guidelines and use the term small EVs (sEVs), rather than exosomes, as the overlapping in size, similar morphology and variable composition make the distinction of EVs type by current purification methods difficult (Lötvall et al. 2014; Théry et al. 2018).

Interestingly, EVs are produced during external exposure to X-rays as part of many cellular processes, including activation of the immune system (Al-Mayah et al. 2015; Jelonek et al. 2016; Diamond et al. 2018). However, to our knowledge, no study has assessed their role during Auger radioimmunotherapy (RIT). Here, we report preliminary results on the role of sEVs as mediators of bystander and systemic effects in melanoma cells exposed to Auger RIT or X-rays.

Materials and methods

Cell lines, antibody and radiolabeling, X-rays

B16F10 mouse melanoma cells (from ATCC) were grown in RPMI1640 medium supplemented with 10% fetal bovine serum (FBS) and 1% penicillin. For exosome collection, cells were incubated in Opti-MEM medium (31985070, Thermo Fisher Scientific), to avoid contamination by fetal bovine serum EVs. B16F10 cells were cultured at 37°C and 5% CO\(_2\), and tested regularly for mycoplasma contamination and for rodent pathogens. B16F10 cells were chosen because they are a well-established and widely used model of melanoma that allows investigating the immune system involvement in vivo (Overwijk and Restifo 2000).

The anti-mouse tyrosinase-related protein 1 (TYRP-1, TRP-1) TA99 mAb (IgG2a format) was purchased from BioXcell. TYRP-1 is the most abundant glycoprotein synthesized by melanocytes and pigmented melanoma cells, and is specific to melanocytes, and primary and metastatic melanomas (Théy et al. 2017). For Auger RIT, TA99 radiolabeling was performed according to the IODO-GEN (1,3,4,6-tetrachloro-3a, 6a-diphenylglycoluryl) (Sigma-Aldrich, St. Louis, MO) method (\(^{125}\)I-TA99) at the specific activity of 370 MBq/mg as previously described (Paillas et al. 2016). TA99 antibody radiolabeling was performed according to the IODO-GEN (1,3,4,6-tetrachloro-3a, 6a-diphenylglycoluryl) (Sigma-Aldrich, St. Louis, MO) method (Pouget et al. 2008). \(^{125}\)I-TA99 specific activity was 370 MBq/mg of protein, and immunoreactivity was 60–95%. All experiments used \(^{125}\)I-TA99 activities ≤4 MBq/mL. Cells were also exposed to X-rays (XenX\(^\circ\), Xstrahl, 225 kV; 2 Gy/min).

Preparation of conditioned medium from X-ray- and RIT-treated cells

The protocol shown in Figure 1(A) was followed. Typically, 50–300 B16F10 donor cells/well were seeded in 6-well plates containing 2 mL of medium. The following day, cells were exposed to \(^{125}\)I-TA99 (0–4 MBq/mL) (RIT) for 48 h or to 0.5 Gy X-rays. The dose of 0.5 Gy X-rays was chosen as bystander effects have been shown to occur at dose below 1 Gy (Nagasawa and Little 1992; Belyakov et al. 2001; Prise and O’Sullivan 2009). Then, cells were washed with Dulbecco’s phosphate-buffered saline (PBS) twice before addition of fresh Opti-MEM\(^\circ\) medium (for sEVs collection) or RPMI 1640 medium with 10% FBS (for bystander experiments) for 2 h. The choice of 2 h incubation was based on previous experiments using other cell lines showing that conditioned medium (CM) from Auger RIT-treated cells was cytotoxic and genotoxic (Paillas et al. 2013, 2016). Then, CM was transferred to recipient cells for bystander effect measurement or used for sEVs isolation.

Determination of direct and bystander cytotoxicity induced by X-rays and Auger RIT

CM was added to untreated recipient B16F10 cells to determine the bystander effects, while fresh non-radioactive medium was added to donor cells after RIT/X-ray exposure to determine the direct effects. Colonies were counted after 10 days of culture as described in (Paillas et al. 2013) to determine clonogenic survival.

In vitro generation of small extracellular vesicles (sEVs)

sEVs were purified from CM samples at 4°C with the protocol described by (Théry et al. 2006) (Figure 1(A)). First, CM samples were centrifuged at 300 x g for 5 min and then at 2000 x g, for 20 min to discard living and dead cells,
respectively, contained in the pellet. Supernatants were then ultracentrifuged at 10,000 × g for 30 min to discard largest vesicles, and the supernatant was then ultracentrifugated at 100,000 × g for 3 h to pellet sEVs fraction. The supernatant was discarded and sEVs were resuspended with PBS, filtered through 0.22 μm filters before a second ultracentrifugation time under the same conditions. Finally, sEVs-containing pellets were resuspended in 200–600 μL of PBS/25 mM trehalose and stored at −80°C (following Le Saux et al. 2020). Cells were counted following the CM collection.

Figure 1. (A) B16F10 cells were exposed to 4 MBq/mL 125I-TA99 mAb for 2 days or to X-rays (0.5 Gy). Immediately after treatment, cells were washed with PBS twice, and Opti-MEM medium was added for 2 h or 48 h in one experiment for the injection of mice with X-rays sEVs (panel 1). The resulting conditioned medium containing sEVs (panel 2) was ultracentrifuged to discard cells, debris, microvesicles (panel 3). Supernatant was ultracentrifugated twice and the pellet containing sEVs was stored at −80°C in PBS/trehalose before NTA (panel 4). (B) Clonogenic survival was assessed in donor and recipients B16F10 cells 10 days after exposure to 125I-TA99 mAb (from 0 to 4 MBq/mL) for two days (donor cells), or incubation with CM in which donor cells were cultured for 2 h (recipient cells). (C) Clonogenic survival was assessed in donor and recipient cells 10 days after exposure to X-rays (0–2 Gy) (donor cells) or incubation with CM in which donor cells were cultured for 2 h (recipient cells). Results are the mean ± SD of three experiments performed in triplicate. *p < .05, **p < .01, ***p < .001, ****p < .0001.
**Extracellular vesicles characterization**

sEVs concentration (number of particles per mL) and size were determined by nanoparticle tracking analysis (NTA) using a NanoSight® NS300 instrument (Malvern Panalytical, Malvern, UK). Suspensions were diluted with particle-free PBS to obtain a particle concentration in the $1 \times 10^4$–$1 \times 10^9$ particles/mL range, as recommended by the manufacturer. Measurements were performed with a 405 nm laser. sEVs suspensions were analyzed using the NanoSight NTA 3.2 software following a tailored script: temperature was set at 25°C, syringe pump at 40 AU (arbitrary unit), three videos of 60 s were recorded. Videos were recorded with a camera level set to 16 and analyzed with a detection threshold set to 6. The number of sEVs released by one cell can be estimated by dividing the total number of particles measured by NTA by the total number of cells.

The expression of some proteins generally associated with sEVs expressed by endosomal sourced vesicles (exosomes) was determined by western blotting using antibodies against ALIX (E6P9B; rabbit mAb, 92880S, Cell Signaling Tec), ADAM10 (ab1997, Abcam, Paris, France), TSG101 (14497-1-AP, Proteintech), and CD81 (D5O2Q; rabbit mAb, 10037, Cell Signaling Technology Inc, Beverly, MA USA). Briefly, $1 \times 10^6$ EV particles were lysed in RIPA buffer supplemented with 1% NaVO₄, phenylmethylsulfonyl fluoride (PMSF), and Protease Inhibitor Cocktail (sc-24948A, Santa Cruz Biotechnology, Heidelberg, Germany) on ice for 20 min. Then, samples were heated at 95°C for 5 min after addition of 4X Laemmli Sample Buffer (1610747, Bio-Rad, Marnes-la-Coquette, France) with $\beta$-mercaptoethanol (M3148, Sigma-Aldrich, St. Quentin Fallavier, France). Proteins were separated by SDS–PAGE and transferred to nitrocellulose membranes (170-4159, Bio-Rad) that were incubated with primary antibodies (1:1000) overnight. As there are not housekeeping proteins clearly identified so far for sEVs, Western blot analysis aimed at providing qualitative information only. Immune reactions were detected with horseradish peroxidase-conjugated goat anti-rabbit secondary antibodies (1:1000, 1 h) (7074S, Cell Signaling Technology, Danvers, MA, USA) and the enhanced chemiluminescence (ECL) detection system (1705060, Bio-Rad).

**Immunodetection of cytosolic double-stranded DNA (dsDNA) in B16F10 cells**

Based on the protocol by (Spada et al. 2019), B16F10 cells grown in 12 mm diameter coverslips were fixed with 4% PFA and permeabilized for 7 min. After blocking nonspecific binding sites using a 1% bovine serum albumin (BSA)/PBS buffer, cells were incubated with an anti-dsDNA antibody (ab27156, Abcam), diluted to 1:1000 in 1% BSA/PBST, at 4°C overnight. After three washes in 1X PBS, cells were incubated with a goat anti-mouse IgG Alexa Fluor® 488 at room temperature for 1 h. After three washes in 1X PBS, coverslips were mounted on a slide using one drop of Vectashield® antifade mounting medium with DAPI (H-1200, Vector Laboratories). Images were acquired at 40× magnification using a Zeiss® Apotome.2 microscope.

**Quantification of sEVs dsDNA content**

Based on the method by (Spada et al. 2020), sEVs dsDNA was extracted using the QIAamp DNA Mini Kit (51304, Qiagen, Courtaboeuf, France), and its concentration was measured using the fluorescent dsDNA-binding dye QuantiFluo® ONE dsDNA System (Promega Corporation, Madison, WI, USA). The amount of sEVs dsDNA (in fg) was estimated relative to the dsDNA concentration (fg/µL) measured with the standard lambda dsDNA curve and corrected by the dilution factor and the initial volume of purified sEVs per cell.

**Cytotoxicity of Auger RIT- and X-ray-generated sEVs**

About 50–70 B16F10 donor cells/well were seeded in 6-well plates containing 2 mL of medium before incubation with $0–4 \times 10^6$ sEVs/mL isolated from CM samples of Auger RIT- or X-ray-treated cells. Clonogenic assays were performed as before.

**Mice**

Female athymic nude (Crl:NU(NCr)-Foxn1nu) and C57BL/6J mice (6–8-week-old; Charles River) were acclimated for 1 week. They were housed at 22°C and 55% humidity with a light-dark cycle of 12 h, and food and water ad libitum. Body weight was determined weekly, and mice were clinically examined throughout the study. Animal experiments were performed in compliance with the French government guidelines and the INSERM standards for experimental animal studies (agreement B34-172-27). They were approved by the Institut de Recherche Cancérologie de Montpellier (IRCM/INSERM) and the Languedoc Roussillon region (CEEA LR France No. 36) (reference number: 1353-9340) ethics committees for animal experimentation. 5 $\times 10^4$ B16F10 melanoma cells in 100 µL of culture medium (without FBS) were inoculated subcutaneously in mice. Amounts of sEVs intratumorally administered in mouse were based on Matsumoto et al. study (Matsumoto et al. 2017) in which 10 µg exosomes were split in 3 injections. This value was drastically reduced to take into account constraints due to radioactive handling. Finally, when tumors reached 20 mm³, mice (n = 8 mice/group) received four consecutive daily intratumoral injections of $4 \times 10^6$ sEVs (0.01 µg) in 50 µL PBS/trehalose each injection, purified from cells exposed to 0.5 Gy X-rays or to 4 MBq/mL $^{125}$I-anti-TA99, or vehicle alone (PBS/trehalose). Tumor growth was monitored every two days by caliper measurement, and mice were euthanized when the tumor reached a volume of 2000 mm³. Tumor volume was calculated using the following formula: length × width × width/2.

**Statistics**

Data were expressed as mean ± standard deviation (SD) for in vitro experiments, and as mean ± standard error of mean (SEM) for in vivo experiments. For clonogenic survival,
comparison between groups were performed using non-parametric Mann–Whitney t-tests. A p-value ≤ .05 was considered significant. Statistical analyses were done with GraphPad Prism® 8.4.3.

Results

**Bystander effects and EVs production are induced during Auger RIT**

Exposure to \(^{125}\)I-TA99 mAb strongly decreased clonogenic survival of B16F10 melanoma cells in an activity-dependent manner (40 ± 7% at 4 MBq/mL; \(p < .0001\) compared with untreated cells) (Figure 1(B)). Similarly, clonogenic survival of recipient B16F10 cells incubated in CM decreased to 62 ± 9% (\(p < .0001\)). Clonogenic survival of donor cells exposed to 2 Gy X-rays decreased to 68 ± 6% (\(p < .0001\)). Conversely, no bystander cytotoxicity was observed in recipient cells incubated with CM from cells exposed to 2 Gy X-rays (105 ± 9%, Figure 1(C)). We determined in a previous study using a nonspecific \(^{125}\)I- PX mAb that soft X-rays produced by \(^{125}\)I do not contribute to cell killing and that resulting mean nucleus absorbed dose is about 0.2 Gy (Pouget et al. 2008).

To test the role of sEVs in the observed bystander effects, exosomes were purified from CM samples from 536 cells exposed to Auger RIT or X-rays (Figure 2(A)). sEVs 537 number per cell varied between 2 and 56 and between 1 538 and 91 for Auger RIT- and X-ray-treated cultures. Size distribution (NTA) was similar for both conditions, with a 539 mean diameter of 128 ± 12 nm and 122 ± 8 nm, respectively (Figure 2(A)). Western blotting confirmed the presence of 540 the sEVs markers ALIX, ADAM10, TSG101 and CD81 (Figure 2(B)).

**Figure 2.** (A) NTA of sEVs purified from NI (\(n = 3\)), Auger RIT-treated (\(n = 4\)) and X-ray-irradiated (\(n = 3\)) samples to estimate the size distribution, calculated mode (nm), and average number of sEVs released per cell. (B) ALIX, ADAM10, TSG101, CD81 protein expression was determined by western blotting after protein extraction from purified sEVs. (C) Clonogenic survival was assessed in B16F10 cells 10 days after exposure to sEVs purified from CM samples collected from Auger RIT- (left panel) or X-ray-exposed (right panel) donor cells. Experiments were performed three times in triplicates.
X-ray- and Auger RIT-generated sEVs are cytotoxic in vitro

Then, B16F10 cells were incubated with various concentrations of purified sEVs (0–400 × 10^6 particles/mL). Clonogenic survival was significantly decreased in a dose-dependent manner, and dropped to 67 ± 32% (p = 0.047) and 21 ± 29% (p < .0001) with 400 × 10^6 sEVs/mL from CM of X-ray- and Auger RIT-treated cells, respectively (Figure 2(C)).

Enrichment of sEVs in dsDNA

Accumulation of cytosolic dsDNA was shown to be progressive during the first 48 h following cell exposure to Auger RIT (Figure 3(A)). Next, quantification of dsDNA in purified sEVs (Figure 3(B)) indicated that its concentration was slightly increased in sEVs isolated from Auger RIT-treated cells compared with sEVs from non-irradiated cells (NI) (0.55 ± 0.18 versus 0.33 ± 0.16 fg per cell; p = .06). Conversely, it was strongly increased in sEVs from X-ray-irradiated cells (1.23 ± 0.44 fg per cell; p = .0007).

Systemic effects of X-ray- and Auger RIT-generated sEVs

Finally, immunocompetent and immunosuppressed (lacking T-cells) mice harboring B16F10 cell xenografts were treated with sEVs isolated from CM of Auger RIT- and
In both mouse strains, sEVs did not delay tumor growth compared with control (vehicle) (Figure 4(A,B)), although, sEVs from CM of X-ray-exposed cells slightly reduced tumor growth in immunocompetent mice between 9 and 15 days post-treatment (Figure 4(B)). This effect was enhanced by injection of sEVs isolated from CM collected after 48 h incubation with X-ray-irradiated cells (preliminary data, n = 3 mice per group). Corresponding enrichment of sEVs in dsDNA was then about 6.1 ± 0.3 fg/cell. This strong efficacy of intratumoral injection of sEVs collected 48 h after onset of irradiation has also been observed in another RIT model using a different type of radiation (unpublished results).

**Discussion**

SEVs secreted from all cell types that participate in local and distal intercellular communication, and thus are attractive candidates for explaining bystander and systemic cytotoxic effects (Al-Mayah et al. 2015; Jelonek et al. 2016; de Araujo et al. 2018). Here, we investigated whether small sEVs released in the supernatant of murine melanoma B16F10 cells exposed to Auger RIT contributed to the in vitro bystander cytotoxic effects. We also tested their in vivo cytotoxic effects after injection in tumor xenografts in immunosuppressed and immunocompetent mice. To compare our results with previous data from our group, we used the...
same in vitro protocol (Paillas et al. 2013; Piron et al. 2014; Paillas et al. 2016; Ladjouhounlou et al. 2019). Donor cells were exposed to $^{125}$I-TA99 mAb for 48 h, and CM was prepared by incubating donor cells with exosome-free fresh medium for 2 h.

SEVs obtained from CM samples from Auger RIT- and X-ray-exposed cells, were shown to express ALIX, ADAM10, TSG101 and CD81, proteins generally associated with sEVs produced by endosomal sourced vesicles (exosomes). These sEVs caused cytotoxic effects in vitro at a concentration of $4 \times 10^8$ particles/mL. The extrapolation between the concentration of purified sEVs and their concentration in CM samples used for bystander experiments is not straightforward. Indeed, sEVs were purified from large CM volumes through successive ultracentrifugation steps (Figure 1(A)). Therefore, it is likely that their concentration was much higher than in CM samples. This could explain why bystander effects were not observed with CM from cells exposed to 0.5 Gy X-rays (Figure 1(C)), whereas clonogenic survival was significantly reduced by sEVs purified from such CM (Figure 2(C)).

However, sEVs cytotoxicity was not confirmed in mice harboring B16F10 cell xenografts which may suggest that they do not induce bystander or systemic effects in vivo. This could be due to the strong aggressiveness of melanoma tumors, which finally overwhelms the cytotoxicity of 4 injections of sEVs. The aggressiveness of the latter tumors was also observed during Auger and alpha RIT as high activities, as compared with other carcinoma models, were required to delay tumor growth (data not shown). Moreover, we did not check the fate of intratumorally injected sEVs. Matsumoto et al. demonstrated that sEVs administered in B16BL6 tumors remained within the tumor up to 48 h post-injection (Matsumoto et al. 2017). In another study, Smyth et al. found that intravenously-injected vesicles were cleared rapidly while those administered in tumors were still detected at 24 h (Smyth et al. 2015). Conversely, we know that during Auger RIT, radiolabeled antibodies remain bound to tumors for several days and this stimulation could therefore be able to generate continuous induction of sEVs secretion by tumor cells. We assumed that the dsDNA contained in sEVs (Figure 3(B)), known to act as danger damage-associated molecular pattern (DAMP) signals, would be determinant in triggering the bystander and systemic responses. However, it must be kept in mind that bystander and systemic cytotoxic effects of sEVs may involve totally different mechanisms.

The role of dsDNA and its shuttling via exosomes during external radiotherapy has been highlighted in recent studies (Vanponuille-Box et al. 2017; Diamond et al. 2018). Cytosolic dsDNA acts as an activator of the DNA sensor cGAS and of its downstream effector STING that will lead to type I-IFN release and T-cell recruitment and activation and ultimately to systemic tumor rejection.

We hypothesize that dsDNA concentration in sEVs from CM of Auger RIT-exposed cells was not enough to observe an anti-tumor immune response. In agreement, sEVs isolated from CM of X-ray-exposed cells, in which dsDNA concentration was higher, showed a slight tumor growth delay in vivo. This effect was higher with sEVs collected from CM collected after 48 h incubation with X-ray-irradiated cells which was accompanied by a higher dsDNA enrichment. It must be noted that the size ($122 \pm 8$ nm vs $129 \pm 8$ nm) and expression of sEVs biomarkers (TSG101, CD81, ALIX, ADAM10) was not modified when CM incubation time was increased. However the number of collected sEVs was significantly increased ($\times 30 \pm 25$) (data not shown). This suggests that dsDNA abundance in sEVs might be influenced by the incubation duration before CM collection. It must be noted that Auger RIT mechanisms of action strongly differ from that of X-rays. This could explain different yield and kinetic induction of dsDNA between the two forms of irradiation. Indeed, the whole cell and the nucleus are directly irradiated by X-rays in a homogeneous way. Conversely, $^{125}$I-mAbs targeting the cell membrane mostly deliver the dose to the cell membrane (Arnaud et al. 2016). This energy deposit leads to the formation of ceramide-enriched large domain (lipid rafts) that participate in downstream signaling cascades involving p38, JNK, NF-kB and reactive oxygen species production. Therefore, the formation of DNA double-strand breaks (DSBs) and micronuclei (Piron et al. 2014; Paillas et al. 2016), which possibly will contribute to dsDNA presence in sEVs, is delayed compared with X-rays. Another factor modulating DNA DSBs is that during RIT, lesions are repaired at the same time as they are generated (Piron et al. 2014) and it is necessary to wait for the radiation absorbed dose to accumulate to overcome DNA repair. Piron found that DNA DSBs were increased only after 24 h post-Auger RIT initiation in a colorectal cancer model (Piron et al. 2014). Then, unrepaired or misrepaired DNA would lead to the formation of micronuclei, that in turn would release their DNA into cytosol before being incorporated into sEVs (Figure 3(C)) (Hintzsche et al. 2017; Takahashi et al. 2017). Then, if the yield of micronuclei depends on the cell repair capacity, it depends also on the radiation absorbed dose, a parameter that was not determined here and should be further investigated. The absorbed dose can also play a role by modulating the activation of the DNA exonuclease TREX1, leading to the digestion of cytoplasmic dsDNA and subsequent tempering of the immune response. Indeed, Vanponuille-Box et al. (2017) found in TSA cells that after a single dose of X-rays higher than 12 Gy, cytosolic dsDNA was cleared by TREX1, precluding activation of the cGAS pathway to induce IFN-1, and thus abolishing the anti-tumor immune response. Therefore, targeted radionuclide dosimetry is a crucial parameter that needs to be considered.

**Conclusion**

We showed that sEVs isolated from CM of Auger-RIT-exposed cells are associated with about 30–40% bystander cytotoxic effects in vitro. Cytoplasmic dsDNA progressively increased in B16F10 cells during Auger-RIT, and was enriched in sEVs purified from CM samples of exposed cells. However, in vivo use of sEVs for cancer treatment showed no therapeutic effects under our conditions. Based on our X-ray results, it is not possible to exclude that other
conditions (absorbed radiation dose and sEVs collection time) would not modulate differently the production of sEVs differently during RIT Auger together with their cytotoxic effects.

**Disclosure statement**

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