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Abstract

Background: Radiotherapy, administered in fractionated as well as ablative settings, is an essential treatment component for breast cancer. Besides the direct tumor cell death inducing effects, there is growing evidence that immune mechanisms contribute - at least in part - to its therapeutic success. The present study was designed to characterize the type and the extent of cell death induced by fractionated and ablative radiotherapy as well as its impact on the release of monocyte migration stimulating factors by dying breast cancer cells.

Methods: Cell death and senescence assays were employed to characterize the response of a panel of breast cancer cell lines with different receptor and p53 status towards γ-irradiation applied in a fractionated (daily doses of 2 Gy) or ablative setting (single dose of 20 Gy). Cell-free culture supernatants were examined for their monocyte migration stimulating potential in transwell migration and 2D chemotaxis/chemokinesis assays. Irradiation-induced transcriptional responses were analyzed by qRT-PCR, and CD39 surface expression was measured by flow cytometry.

Results: Fast proliferating, hormone receptor negative breast cancer cell lines with defective p53 predominantly underwent primary necrosis in response to γ-irradiation when applied at a single, ablative dose of 20 Gy, whereas hormone receptor positive, p53 wildtype cells revealed a combination of apoptosis, primary, and secondary (post-apoptotic) necrosis. During necrosis the dying tumor cells released apyrase-sensitive nucleotides, which effectively stimulated monocyte migration and chemokinesis. In hormone receptor positive cells with functional p53 this was hampered by irradiation-induced surface expression of the ectonucleotidase CD39.

Conclusions: Our study shows that ablative radiotherapy potently induces necrosis in fast proliferating, hormone receptor negative breast cancer cell lines with mutant p53, which in turn release monocyte migration and chemokinesis stimulating nucleotides. Future studies have to elucidate, whether these mechanisms might be utilized in order to stimulate intra-tumoral monocyte recruitment and subsequent priming of adaptive anti-tumor immune responses, and which breast cancer subtypes might be best suited for such approaches.

Keywords: Ablative radiotherapy, Fractionated radiotherapy, Monocyte migration, Chemokinesis, Chemotaxis, Nucleotides, Dying cell clearance, Immunogenic cell death, Anti-tumor immunity
Background

Radiotherapy is a crucial treatment part for the management of breast cancer. Commonly, it is applied in daily fractions of 1.8-2 Gy for 5 to 7 weeks to a total dose of 50 to 66 Gy [1-5]. Fractionated irradiation schemes are considered to be beneficial for the reduction of tumor burden, since they are intended to exploit the discrepant DNA repair capacity of tumor and normal tissue. Larger, irreparable damage per unit dose is induced in the tumor, whereas efficient DNA repair mechanisms compensate for the damage induced by the daily irradiation fractions in the adjacent normal tissue. Moreover, tumor reoxygenation can occur between the fractions and tumor cells can redistribute to more radiosensitive phases of the cell cycle [6]. At present several clinical trials support the use of higher doses per fraction in order to shorten the overall treatment period. In the UK doses of 2.66 Gy per fraction are already accepted practice [7,8], and ongoing trials will determine in how far even higher single doses (5.7-6 Gy) may define a new optimum. Additionally, there are specific applications, where ablative, large single doses of 10–25 Gy are delivered locally to the tumor, for instance during intra-operative radiotherapy (IORT) [9-11].

The tumor cell death inducing effect of radiotherapy has been regarded as the major determinant of its therapeutic success for a long time. Nevertheless, there is accumulating experimental evidence that innate as well as adaptive immune responses contribute - at least in part - to the reduction of tumor burden and tumor control [12]. Essentially, previous studies have revealed that radiotherapy stimulates a type I interferon-dependent priming of adaptive anti-tumor immune responses, including tumor-specific CD8+ cytotoxic T cells, by antigen-presenting cells (APCs) [13-17]. These effects were only observed in case of ablative but not fractionated radiotherapy, and the underlying mechanisms remain largely elusive. Conceivably, tumor cells respond differently to γ-irradiation when applied in an ablative or a fractionated setting. The mode as well as the extent of cell death might vary substantially. And since different types of cell death are well known to have different immunological consequences, we supposed that the extent as well as the type of tumor cell death in response to radiotherapy might govern and shape the subsequent anti-tumor immune responses [18]. For systemic anthracycline therapy it has been reported that danger signals released from dying tumor cells trigger the activation of APCs and subsequent T cell priming [19]. One of the initial steps in this scenario was the intra-tumoral recruitment of monocytic precursor cells, which then differentiated into highly potent APCs [20]. Therefore, the present study was designed to analyze the cell death response of different breast cancer cell lines towards fractionated and ablative radiotherapy and its impact on the release of signaling molecules, which stimulate monocyte migration. We observed that fast proliferating, hormone receptor negative, and p53 mutant cell lines predominantly underwent necrosis in response to radiotherapy, particularly when administered in an ablative regime. Necrosis induction was paralleled by the release of apyrase-sensitive nucleotides, which effectively stimulated monocyte migration and chemokinesis. In hormone receptor positive breast cancer cells with functional p53 this process was impaired by irradiation-induced surface expression of the ectonucleotidase CD39 that degrades extracellular nucleotides. Hence, our study opens the question, whether ablative radiotherapy might be utilized for targeted necrosis induction in fast proliferating, hormone receptor negative breast cancer with defective p53 in order to stimulate intra-tumoral monocyte recruitment and subsequent priming of adaptive anti-tumor immune responses.

Methods

Cells and reagents

The human breast cancer cell lines MCF7, BT474, HCC1937, HCC1806, MDA-MB468, and BT549 were obtained from ATCC (Manassas, VA, USA) or CLS (Heidelberg, Germany), and were cultured in RPMI-1640 medium supplemented with 10% heat-inactivated fetal calf serum (FCS), 100 units/ml penicillin, 0.1 mg/ml streptomycin, and 10 mM HEPES (all from Life Technologies, Karlsruhe, Germany) at 37°C and 5% CO₂ (MCF7, BT474, HCC1937, and HCC1806), or in DMEM (BT549), or DMEM/F12 (1:1) medium (MDA-MB468) supplemented with 10% heat-inactivated FCS, 100 units/ml penicillin, and 0.1 mg/ml streptomycin at 37°C and 7.5% CO₂, respectively.

THP-1 cells were obtained from ATCC and were cultured in RPMI-1640 medium supplemented with 10% heat-inactivated FCS, 100 units/ml penicillin, 0.1 mg/ml streptomycin, and 10 mM HEPES. Preparation of human peripheral blood monocytes was carried out as described previously [21]. Briefly, PBMCs were prepared from heparinized blood of healthy volunteers by Biocoll density gradient centrifugation (Biochrom AG, Berlin, Germany). Monocytes were positively selected from PBMCs with anti-CD14 magnetic beads (Miltenyi, Bergisch Gladbach, Germany) according to the manufacturer’s recommendations and allowed to recover for 1 day in X-Vivo 15 medium (Lonza, Basel, Switzerland) supplemented with 10% autologous serum, 100 units/ml penicillin, and 0.1 mg/ml streptomycin before further use.

The p53 status of all breast cancer cell lines was determined by cDNA sequencing. Full length PCR products were generated from cDNA (80 ng per reaction) by employing 5 units HotStar HiFidelity DNA-Polymerase in 1 x HotStar HiFidelity reaction buffer, and 1 x Q solution (all from Qiagen, Hilden, Germany) in the presence of 1 μM of each primer (p53 Forward 5′-ATG
GAG GAG CCG CAG TCA G-3', p53 Reverse 5'-TCA GTC TGA TGC AGG CCC TTC T-3', synthesized by Sigma-Aldrich, Taufkirchen Germany) in 100 µl final volume (cycling program: 1 x 5' 95°C; 40 x 15' 95°C, 1' 60°C, 1' 30'' 72°C; 1 x 10' 72°C). The amplicons were purified by the NucleoSpin Extract II Kit (Macherey & Nagel, Düren, Germany), and sequencing was performed by Seqlab Sequencing Services (Goettingen, Germany).

Carboxbenzoxyl-valyl-alanyl-aspartyl-[O-methyl]-fluoromethylketone (zVAD-fmk) was obtained from Bachem (Bubendorf, Switzerland), necrostatin-1 from Enzo Life Sciences (Loerrach, Germany), calcine-AM from Merck Calbiochem (Darmstadt, Germany), bafilomycin A1 and ARL-67156 from Tocris R&D Systems (Wiesbaden, Germany). The annexin V-FITC apoptosis detection kit, anti-CD39-PE, anti-CD73-FITC, and anti-CD203c-APC antibodies were purchased from BD Biosciences (Heidelberg, Germany), and 5-dodecanoylaminofluorescein-di-β-galactopyranoside (C12-FDG-FITC) was from Life Technologies.

Mouse monoclonal western blot antibodies anti-p21WAF1 and anti-vinculin were obtained from BD Biosciences or Sigma-Aldrich, respectively. The chemokines SDF-1α and WKYMvm (agonist of formyl-peptide receptors 1, 2, and 3) were from R&D Systems, adenosine 5'-triphosphate disodium salt (ATP) from Sigma-Aldrich, and nucleotide diphosphohydrolase (apyrase) was purchased from New England Biolabs (Frankfurt, Germany).

Growth analyses and determination of doubling times
Cells were seeded into 24-well plates (2.5 x 10⁴ cells per well) and allowed to adhere for 5 h. Medium was replaced (medium supplemented with 10% or 2.5% FCS was used as indicated), and cells were grown for up to 4 days. Every day, cells were harvested by trypsinization and cell numbers were determined by counting. Growth curves were generated by plotting log cell number (y-axis) versus time (x-axis), and doubling times were calculated on the basis of the slopes of the corresponding regression lines.

X-ray treatment and production of cell-free culture supernatants
Cells were seeded into 6-well (0.5-1 x 10⁶ cells per well) or 24-well plates (0.25-1 x 10⁵ cells per well) in culture medium supplemented with 10% FCS and allowed to adhere overnight. Immediately prior to irradiation, culture medium was replaced by serum-reduced medium (2.5% FCS). Cells were irradiated at the indicated doses with a Mueller RT-250 γ-ray tube (200 kV and 10 mA, Thoraeus filter, 1 Gy in 1 min 52 s). Fractionated irradiation was carried out daily. Cell-free supernatants were collected by centrifugation (10,000 g, 5 min, 4°C) at the indicated time points and stored at −80°C until further use.

SDS-PAGE and Western blot analyses
Reducing 6-15% gradient SDS-PAGE and Western blot analyses of whole cell lysates were performed as described previously [22,23] with 300 µg protein extract per lane. After electrophoretic separation, proteins were transferred to PVDF Immobilon FL membranes (Merck Millipore, Darmstadt, Germany). Membranes were blocked with 5% low-fat milk in TBST buffer (13 mM Tris–HCl pH 7.5, 150 mM NaCl, and 0.02% Triton X-100) and incubated with monoclonal mouse antibodies against p21WAF1 (BD Biosciences) or vinculin (Sigma-Aldrich), respectively. After incubation with the corresponding IRDye-conjugated secondary antibodies (LI-COR Biosciences, Bad Homburg, Germany) and extensive washing in TBST buffer, IRDye fluorescence was read with a LI-COR Odyssey scanner.

Flow cytometric measurement of phosphatidylserine externalization, plasma membrane integrity, senescence-associated β-galactosidase activity, and ectonucleotidase surface expression
All FACS measurements were performed on an LSRII cytometer (BD Biosciences) and data were analyzed with FACSDiva (BD Biosciences) or FlowJo 7.6.3 Software (Tree Star Inc., Ashland, OR, USA), respectively.

Externalization of phosphatidylserine (PS) and plasma membrane integrity were measured by staining with annexin V-FITC/propidium iodide (annexin V staining kit, BD Biosciences) as described previously [24]. Briefly, 1 x 10⁵ cells were incubated with 5 µl FITC-labeled annexin V in 50 µl staining buffer (both from BD Biosciences) supplemented with 5 µg/ml propidium iodide (PI, Sigma-Aldrich) for 30 min on ice. Following an additional washing step in staining buffer, annexin V-FITC and PI fluorescence were assessed by flow cytometry. Cells with positive annexin V-FITC but negative PI signal were considered apoptotic, and cells double-positive for annexin V-FITC and PI staining were considered necrotic. In order to distinguish primary from secondary necrotic cells, the poly-caspase inhibitor carbobenzoxy-valyl-alanyl-aspartyl-[O-methyl]-fluoromethylketone (zVAD-fmk, Bachem) was employed. zVAD-fmk blocks apoptosis and the transition into secondary necrosis. Hence, annexin V-FITC/PI double positive cells, which were detected in the presence of zVAD-fmk, were considered primary necrotic.

Senescent cells were stained by using 5-dodecanoylaminofluorescein-di-β-galactopyranoside (C12-FDG-FITC) or 5-dodecanoylaminofluorescein-di-β-galactosidase [25]. At the indicated time points after irradiation, cells were incubated with 100 nM bafilomycin A1 (Tocris R&D Systems) in serum-free medium for 1 h at 37°C for lysosomal alkalinization. Subsequently, C12-FDG-FITC was added at a final concentration of 50 µM, and cells were incubated for 1 h at 37°C to
allow substrate conversion. After two washing steps in PBS, cells were collected by trypsinization and analyzed by flow cytometry. Cells with high C12-FDG-FITC and high SSC signal were considered senescent.

For ectonucleotide surface staining, 1 × 10^5 cells were incubated with 2 μl anti-CD39-PE, anti-CD73-FITC, or anti-CD203c-APC in 50 μl FACS staining buffer (all from BD Biosciences) for 30 min on ice. After two washing steps in FACS staining buffer, cells were analyzed by flow cytometry. Relative surface expression was calculated as the median fluorescence intensities of anti-ectonucleotide staining subtracted by the corresponding isotype controls.

**Transwell migration assay**

Transmigration assays were performed in 96-well Multiscreen-MIC transwell chambers with 5 μm pore size (Merck Millipore) as described before [22,23]. In brief, 1 × 10^5 calcein-labeled THP-1 cells per well were seeded in a final volume of 80 μl onto the 96-well filter Plate. 320 μl of supernatants or chemokines dissolved in serum-free RPMI-1640 medium were added to the lower chamber. The filter was mounted onto the lower chamber and transmigration was allowed for 90 min at 37°C. Subsequently, the cells in the lower chamber were collected by centrifugation and lysed in 100 μl lysis buffer (20 mM HEPES-K pH 7.4, 84 mM KCl, 10 mM MgCl2, 0.2 mM EDTA, 0.2 mM EGTA, 0.5% Igepal). Green calcein fluorescence was quantified with a Synergy MX fluorescence reader (BioTek Instruments GmbH, Bad Friedrichshall, Germany), and transmigration was calculated as percentage of total cells deployed.

In some experiments, supernatants were subjected to ultrafiltration with VivaSpin 2 centrifuge tubes with an exclusion limit of 10 kDa (Sartorius Stedim Biotech, Goettingen, Germany) as described before [22]. After passing the entire liquid phase through the filter, the filter was rinsed well with culture medium and the volume of the two fractions (substances smaller and larger than 10 kDa) was readjusted to the initial volume employed. Then, the fractions were applied to a transmigration assay.

Apyrase treatment was carried out by adding 500 milliunits of nucleotide diphosphohydrolase (apyrase, New England Biolabs) to 1.5 ml culture supernatant and allowing nucleotide degradation for 30–50 min at 37°C. Heat-inactivated apyrase served as a control. After digestion, supernatants were used in transwell migration or chemotaxis/chemokinesis assays, respectively.

**Chemotaxis/chemokinesis assay in IBIDI μ-slide 2D chemotaxis chambers**

IBIDI μ-slide 2D chemotaxis chambers (IBIDI, Munich, Germany) were used to analyze chemotaxis and/or chemokinesis of primary human monocytes by live cell tracking as described before [22]. Briefly, monocytes were seeded into the observation area of the chamber in X-Vivo 15 medium supplemented with 5% autologous serum. Adherence was allowed for 15 min, and non-adherent cells were carefully washed away. The reservoirs of the chamber were filled with medium supplemented with 5% autologous serum, and the stimulus was added to the upper reservoir. The slide was mounted onto the heated stage of an AxioObserver Z1 inverted microscope (Zeiss, Goettingen, Germany), and time-lapse video microscopy was performed at 37°C and 5% CO2 for 3 h at 5 × magnification. Pictures were taken every 2 min and migration of 40 randomly picked cells was tracked with the ImageJ manual tracking plug-in. Accumulated distance, euclidean distance (linear distance between start and end position), and y forward migration index (yFMI = mean of (endpoint in y direction/accumulated distance) of all cells analyzed) were determined with the IBIDI chemotaxis and migration tool (IBIDI). Analysis window was set from 10 min to 2 h and 10 min (2 h time frame). Apyrase treatment of culture supernatants was performed as described for transwell migration assays.

**Quantitative Realtime RT-PCR**

RNA isolation and qRT-PCR analyses were performed as described previously [26,27]. Briefly, total RNA was extracted with the NucleoSpin RNA II Kit (Macherey & Nagel). 1 μg of isolated RNA was subjected to reverse transcription with 200 units RevertAid reverse transcriptase in the presence of 50 μM random hexamers, 5 μM Oligo(dT)18, 400 μM dNTPs, and 1.6 units/μl Ribolock RNase inhibitor (all from Fermentas, St. Leon-Rot, Germany). The resulting cDNA (20 ng per reaction) was applied to qRT-PCR analyses (20 μl final volume) with 300 nM primers (synthesized by Sigma-Aldrich) in 1× Maxima SYBR Green qPCR Mastermix (Fermentas) and a standard cycling protocol (10 min 95°C, 45× (15 s 95°C, 30 s 60°C)) on an LC480 qPCR cycler (Roche Applied Science, Penzberg, Germany). The following primer pairs were used: p21^WAF1 Forward 5′-CTG GAG ACT CTC AGG GTC GAA A-3′, p21^WAF1 Reverse 5′-AGT GGT AGA AAT CTG TCA TGG T-3′, Egr-1 Forward 5′-GAG CAC CTTG ACC GCA GAG TC-3′, Egr-1 Reverse 5′-CCA GCA CCT TCT CTG TGT TCA-3′, 18S rRNA Forward 5′-CGG CTA CCA CAT CCA AAG AA-3′, 18S rRNA Reverse 5′-GCT GGA ATT ACC GCG GCT-3′, β2-microglobulin Forward 5′-TGC TCG CGC TAC TCT CTC TTT C-3′, β2-microglobulin Reverse 5′-TCT CTG CTG GAT GAC GTG AGT AAA C-3′. Relative quantification was performed by employing the standard curve method, and the results were normalized on the means of 18S rRNA and β2-microglobulin. Untreated control cells were used as calibrator.
Results and discussion

A few previous studies have shown that radiotherapy can stimulate anti-tumor immune reactions, which contribute to the reduction of tumor burden [13-17]. In principle, the authors observed a type I interferon-dependent, APC-mediated priming of tumor-specific CD8+ T cell responses. Notably, the induction of these T cell responses was limited to ablative radiotherapy regimes, where γ-irradiation was applied at high single doses of more than 10 Gy. The tumor cell response towards low or high single doses as well as fractionated γ-irradiation in terms of apoptosis, necrosis, and senescence induction is likely to diverge substantially, and different types of tumor cell death are well known to stimulate different types of immunological consequences. Therefore, we hypothesized that the type of tumor cell response, which is induced by different regimes of radiotherapy, might shape the immunological consequences [18,28]. In order to address this issue, we analyzed the tumor cell response towards different regimes of γ-irradiation in three breast cancer lines: HCC1937, MCF7, and BT474. We deliberately chose cell lines of divergent molecular breast cancer subtypes as well as divergent estrogen, progesterone, and Her2/neu receptor status (Figure 1A) [29,30]. The cell lines exhibited clear differences in proliferation rates with short (MCF7, 24 h), intermediate (HCC1937, 46 h), and long (BT474, 77 h) doubling times in the presence of 2.5% FCS (Figure 1B, C). p53 mutation status and function were confirmed by cDNA sequencing and by immunoblot analysis of p21\textsuperscript{WAF1} induction in response to γ-irradiation at 4 Gy. Only MCF7 cells, which have been reported to possess wildtype p53 [30], revealed irradiation-induced upregulation of p21\textsuperscript{WAF1} protein expression starting approximately 4 h and reaching a plateau around 8 h after irradiation.

| Breast cancer cell line | Mol. subtype | Tumor type | ER | PR | Her2/neu | p53 |
|-------------------------|--------------|------------|----|----|---------|-----|
| BT474                   | Luminal      | invas. duct. ca. | +  | +  | +       | 285 E-K |
| MCF7                    | Luminal      | invas. duct. ca. | +  | +  | -       | WT  |
| HCC1937                 | Basal A      | prim. duct. ca. | -  | -  | -       | 306 R=Stop |

Figure 1 Breast cancer cell lines of different origin reveal different doubling times and p53 functionality. (A) Breast cancer cell lines used in the present study. Molecular subtype, tumor type and receptor status have previously been reported [29], and p53 mutational status was determined by cDNA sequencing. (B) Growth curves of breast cancer cell lines were generated in the presence of 2.5% or 10% FCS, respectively. Means ± s.d. of triplicates are shown. (C) Doubling times of exponentially growing cells were calculated from the data shown in (B). (D) p21\textsuperscript{WAF1} induction as an indicator of p53 functionality was examined in whole cell lysates after irradiation with 4 Gy at the indicated time points by 6-15% SDS-PAGE (300 μg protein extract per lane) and subsequent immunoblot analysis. Vinculin served as a loading control.
Different γ-irradiation regimes induce different modalities of cell death and senescence in breast cancer cell lines

Next, we investigated the type of tumor cell response towards different regimes of γ-irradiation. Cells were irradiated at single doses of 2 Gy or 20 Gy, or daily fractions of 2 Gy, respectively, and the percentage of apoptotic, necrotic, and senescent cells was measured by flow cytometry over a period of 4 days after irradiation (Figure 2A, D). In order to distinguish between primary and secondary (post-apoptotic) necrosis, we employed the poly-caspase inhibitor zVAD-fmk, which blocks apoptosis and the subsequent transit into secondary necrosis (Figure 2B). The necroptosis inhibitor necrostatin-1 was used to assess the contribution of necroptosis in our experimental system (Figure 2C). We made the following observations: (i) The strongest response of apoptosis, necrosis, and senescence induction was detected in fast proliferating MCF7 and HCC1937 cells. Slowly proliferating BT474 cells revealed only a moderate degree of apoptosis, necrosis, or senescence induction, respectively. (ii) Fast proliferating, p53 wildtype MCF7 cells underwent a combination of apoptosis and necrosis – primary as well as secondary (post-apoptotic) necrosis. Appearance of senescent MCF7 cells was only observed in response to ablative irradiation with 20 Gy, (iii) HCC1937 cells with mutant p53 predominantly underwent primary, apoptosis-independent necrosis and senescence. (iv) Of all γ-irradiation regimes applied, ablative irradiation at 20 Gy provoked the most pronounced cellular responses in terms of apoptosis, necrosis, and senescence induction. These findings allow the conclusion

**Figure 2** Different γ-irradiation regimes induce different modalities of cell death and senescence in breast cancer cell lines. (A) Induction of apoptosis, necrosis, and senescence. Breast cancer cell lines were left untreated or γ-irradiated at single doses of 2 Gy, 20 Gy, or daily fractions of 2 Gy, respectively. Induction of apoptosis and necrosis was determined 1–4 days after irradiation by annexin V-FITC/PI staining and FACS analysis. Annexin V-FITC positive, PI negative cells were considered apoptotic, double positive cells were considered necrotic. Senescence induction was measured by flow cytometric SA-β-gal staining with the fluorogenic substrate C12-FDG-FITC. Cells with high C12-FDG-FITC and high SSC signal were considered senescent. Means ± s.d. of triplicates are depicted. (B) Induction of apoptosis and necrosis in the presence of zVAD-fmk. PS externalization and plasma membrane integrity were measured as in (A) in the presence of 50 μM of the poly-caspase inhibitor zVAD-fmk. Means ± s.d. of triplicates are shown. (C) Induction of apoptosis and necrosis in the presence of necrostatin-1. PS externalization and plasma membrane integrity were measured as in (A) in the presence of 50 μM of the necroptosis inhibitor necrostatin-1. Means ± s.d. of triplicates are shown. (D) Representative dot plots of HCC1937 cells stained for SA-β-gal activity with the fluorogenic substrate C12-FDG-FITC.
that ionizing irradiation - particularly when applied in an ablative setting of high single doses - primarily induces cell death and senescence in fast proliferating cancer cells. In cells with wildtype p53, a combination of apoptosis, primary, and secondary (post-apoptotic) necrosis is the major consequence, whereas cells lacking functional p53 mainly undergo primary necrosis and senescence. The contribution of necroptosis in this context is only a marginal one. These findings are in line with reports showing that p53 is crucial for irradiation-induced apoptosis, either via transcriptional activation of pro-apoptotic p53 target genes, including Bax, Puma, and Noxa, or via transcription-independent pathways, respectively [31-35]. In addition, p53 has been reported to be essential for establishing and maintaining certain types of senescence [36,37]. Nevertheless, p53-independent forms of senescence apparently do exist [38].

Ablative γ-irradiation induces the release of low molecular weight, apyrase-sensitive factors, which stimulate monocyte chemokinesis

In order to stimulate productive, tumor-specific immune responses by radiotherapy, irradiated, dying tumor cells have to be detected and engulfed by APCs, which subsequently migrate into the draining lymph nodes, process and (cross-) present ingested tumor-antigens, and thus subsequently migrate into the draining lymph nodes, process and (cross-) present ingested tumor-antigens, and thus finally prime adaptive anti-tumor immune responses, in-and (cross-) presenting ingested tumor-antigens, which might contribute to monocyte migration by irradiated, necrotically dying HCC1937 cells.

Different necrotic cell-derived danger signals have been reported to be involved in monocyte recruitment. High molecular weight compounds, like heat shock proteins, high-mobility group box 1 protein (HMGB-1), S100 protein family members, small nuclear ribonucleoproteins, monosodium urate crystals, or nucleic acids, as well as low molecular weight compounds, like nucleotides, have been described [41]. In order to elucidate, which of these factors might contribute to monocyte attraction by ablatively irradiated HCC1937 cells, cell-free supernatants were subjected to ultrafiltration with an exclusion limit of 10 kDa. THP-1 cell migration towards the filtered supernatants was virtually not affected after high molecular weight compounds had been removed. Comparable results were obtained for purified ATP (MW = 507 Da), whereas the classical CXC chemokine SDF-1α (MW = 11 kDa) was more or less completely retained in the fraction with a molecular weight of more than 10 kDa, thus confirming a proof-of-principal of this procedure (Figure 3B left panel). Furthermore, incubation with active but not heat-inactivated nucleotide diphosphohydrolase (apyrase) completely abrogated THP-1 cell migration towards supernatants of HCC1937 cells that had been irradiated at 20 Gy. Again, parallel results were observed for purified ATP, whereas migration towards SDF-1α was basically not impaired by apyrase digestion (Figure 3B right panel). These findings allow the conclusion that the THP-1 cell migration stimulating factors, which are released by ablatively irradiated, necrotically dying HCC1937, are of low molecular weight and sensitive to apyrase treatment, apparently nucleotides. Different studies have previously provided evidence for the involvement of extracellular nucleotides in the recruitment of monocytes, macrophages, and dendritic cells by dying cells in vitro and in vivo [20,42,43]. However, currently it is being controversially discussed, whether nucleotides per se do stimulate directional chemotactic responses in monocytes and macrophages, or if they rather act as auto- and paracrine amplifiers of other chemotactic stimuli, such as complement C5a [44,45].

Therefore, we next characterized the migratory response of primary human monocytes towards supernatants of γ-irradiated HCC1937 cells by time-lapse video microscopy in 2D chemotaxis/chemokinesis chambers. The obtained trajectory paths and the detailed analysis of accumulated and euclidean distance as well as the
forward migration index in direction of the gradient (yFMI) clearly show that the migratory response of primary monocytes towards supernatants of ablatively irradiated HCC1937 cells was a chemokinetic and not a chemotactic one (Figure 4A, B). In comparison to the untreated controls, supernatants of HCC1937 cells irradiated at 20 Gy intensified and accelerated monocyte migration as revealed by significant increases in the accumulated as well as the euclidean distance. However, monocyte migration was not directed towards the chamber, in which the supernatant was applied, and the yFMI value was even negative. Interestingly, parallel results were obtained for purified ATP supporting the conclusion that in our system nucleotides do not stimulate chemotaxis but rather chemokinesis as has already been described by others [44]. For comparison, the chemotactic FPR agonist WKYMVm was employed. Here, the increase in the accumulated distance was comparable to the one obtained with supernatants of ablatively irradiated HCC1937 cells and ATP, but the acquired euclidean distance was noticeably higher, and the yFMI was clearly positive, as the vast majority of cells migrated in the direction of the gradient. Notably, supernatants of HCC1937 cells that had been subjected to the fractionated irradiation scheme with daily doses of 2 Gy stimulated monocyte chemokinesis to a lesser, yet in terms of the accumulated distance still significant extent. Chemokinesis in the presence of supernatants collected from HCC1937 irradiated at a single dose of 2 Gy did not differ from the untreated control. Again, apyrase treatment significantly reduced monocyte chemokinesis stimulated by supernatants of ablatively irradiated HCC1937 cells, and the median accumulated distance declined to the level that was observed with supernatants of viable control cells (Figure 4B, C, D).

Hence, our migration data clearly show that necrotically dying, ablatively irradiated HCC1937 cells - and to a lesser extent also cells subjected to fractionated irradiation with daily doses of 2 Gy - release low molecular weight, apyrase-sensitive nucleotides, which stimulate monocyte chemokinesis in a similar fashion as ATP.

**Ablative γ-irradiation induces the upregulation of CD39 surface expression in MCF7 breast cancer cells**

In contrast to HCC1937 cells, supernatants of MCF7 cells did not stimulate monocyte migration, although MCF7 cells strongly underwent primary and secondary necrosis in response to ablative γ-irradiation with 20 Gy (Figure 2). Complex nucleotide secretion processes, like the caspase/pannexin axis, which is activated during apoptosis and has been described to be impaired in MCF7 cells [46], apparently are of minor importance in case of necrosis-associated nucleotide release, since during necrosis the plasma membrane disintegrates and intracellular contents can passively leak out. Hence, the question that arises is: why do supernatants of necrotic MCF7 cells not contain monocyte migration stimulating nucleotides? A conceivable explanation for this observation would be that MCF7 cells express ectonucleotidases, which degrade extracellular nucleotides [47]. Surface staining of ectonucleotidases revealed that this in fact was the case (Figure 5A, B and data not shown). In contrast to HCC1937 and BT474 cells, MCF7 cells showed a low, but well detectable basal expression of the ectonucleotidase CD39, which was
strongly increased in response to irradiation with 20 Gy and to a lesser extent also by fractionated irradiation with daily doses of 2 Gy (Figure 5A, B). Importantly, pharmacological inhibition of CD39 ectonucleotidase activity by addition of ARL-67156 resulted in the release of comparable amounts of THP-1 cell migration stimulating factors by ablatively irradiated MCF7 cells as had been observed with HCC1937 cells (Figure 5C and Figure 3A). Thus, upregulated CD39 apparently degrades extracellular nucleotides released by necrotically dying MCF7 cells.

The irradiation-induced increase in CD39 surface expression revealed a biphasic kinetics with an initial rise between days 1 and 2 after irradiation and an even stronger increase starting on day 3. The basal expression...
of CD39 in MCF7 cells has already been reported by others, but the mechanisms, which account for the differences in CD39 expression compared to HCC1937 and BT474 cells, are poorly understood [48]. Candidate transcriptional regulators in this regard are p53 and the nuclear hormone receptors for estrogen (ER) and
progesterone (PR), since the three breast cancer lines differ in p53 functionality and hormone receptor status (Figure 1A). In silico analysis of the CD39 promoter region employing the AliBaba 2.1 platform (http://www.gene-regulation.com/pub/programs/alibaba2/index.html) revealed several transcription factor binding sites, including sites for the estrogen receptor (ER) and the progesterone receptor (PR) but no p53 response element (Figure 5D). Yet, p53- and ER-mediated transcriptional regulation appear to be closely interconnected, since they do not only mutually regulate each other’s expression but also have been described to control target gene expression in a coordinate manner [49-52]. Hence, p53 and ER might orchestrate basal CD39 expression in MCF7 cells. Following γ-irradiation, particularly when applied in an ablative scheme, MCF7 cells showed a robust activation of p53 as revealed by induction of p21WAF1 mRNA and protein expression (Figure 5E, Figure 1D). Hence, activated p53 (in cooperation with ER) might account for the upregulation of CD39 expression, since it was only observed in MCF7 cells and the induction of the prototypical p53 target p21WAF1 displayed a comparable biphasic time course as that of CD39. Nevertheless, indirect mechanisms, including the p53-mediated activation of other transcriptional regulators, could also be involved. As such Egr-1, an immediate early response transcription factor, which is well known to be induced and activated by ionizing irradiation and whose response element was identified close to the transcription start site within the CD39 promoter (Figure 5D), was induced in γ-irradiated MCF7 cells in a similar fashion as p21WAF1 and CD39 [53] (Figure 5E). Interestingly, Egr-1 has been reported to interact with p53 and to enhance transcriptional activation by p53 [54,55]. Our data do not allow detailed conclusions on the mechanisms, which govern irradiation-induced upregulation of CD39 expression in MCF7 cells. Nevertheless, they support a scenario, in which p53, ER and Egr-1 could play a crucial role. Further studies are required to elucidate this issue in greater depth and to find out if other transcription factors, such as Sp-1, Stat-3 or NF-kB, also are involved. It should be noted that we also measured the surface expression levels of CD73 and CD203c, two other well-known ectonucleotidases, but we did not detect any basal expression, nor an irradiation-induced upregulation in the three breast cancer cell lines tested (data not shown).

**Fast proliferating breast cancer cells with mutant p53 and a strong necrosis response towards ablative γ-irradiation release factors that stimulate monocyte migration**

In order to examine, whether our findings that fast proliferating, hormone receptor negative HCC1937 breast cancer cells with defective p53 and a robust necrotic response towards ablative γ-irradiation release monocyte migration stimulating nucleotides, are of broader relevance, we analyzed three more cell lines: HCC1806, MDA-MB468, and BT549 cells. These hormone receptor negative breast cancer cell lines with mutant p53 (Figure 6A) exhibited doubling times of 30 h (HCC1806), 51 h (MDA-MB468) or 77 h (BT549), respectively (Figure 6B). Irradiated HCC1806 and MDA-MB468 cells underwent primary necrosis to a strong and comparable degree as HCC1937 cells, whereas in slowly proliferating BT549 cells no significant induction of necrosis was detected (Figure 6C). When combining the results of all cell lines analyzed in the present study, we observed a clear and significant negative correlation between doubling times and necrosis induction by ablative irradiation with 20 Gy. This held true for total as well as primary necrosis in the presence of zVAD-fmk (Figure 6D). Transwell migration assays with THP-1 cells revealed that only supernatants of irradiated HCC1806 and MDA-MB468 but not BT549 cells released monocyte migration stimulating factors. Again, the strongest monocyte migration was observed with supernatants of ablatively irradiated cells (Figure 6E). As expected, the three p53 defective, ER negative cell lines did neither show any basal CD39 surface expression in FACS analyses, nor its irradiation-induced upregulation (data not shown). Finally, we combined the data on THP-1 cell migration and necrosis induction of all p53 mutant, hormone receptor negative breast cancer lines. The cell lines with strong necrosis induction in response to γ-irradiation were the ones, whose supernatants potently stimulated THP-1 cell migration. Pearson correlation analysis revealed a significant positive correlation between the percentage of THP-1 cell migration and the percentage of total necrosis induced. This correlation was even more stringent when only primary necrosis was considered (Figure 6F).

In summary, our study reveals that fast proliferating, hormone receptor negative breast cancer cell lines with defective p53 intensively undergo necrosis in response to γ-irradiation, particularly when applied in an ablative regime at a single dose of 20 Gy. During necrosis, the cells release nucleotides, which efficiently stimulate monocyte migration in a chemokinetic manner. In hormone receptor positive, p53 wildtype cells, like MCF7, this appears to be hampered by irradiation-induced upregulation of CD39, which degrades extracellular nucleotides. Our study opens several questions, including the detailed molecular mechanisms, which orchestrate irradiation-induced upregulation of CD39 and the specific role of p53 and the hormone receptors in this scenario. Moreover, it will be interesting to further characterize the subpopulation of cells, in which CD39 expression is increased in response to irradiation. The biphasic kinetics and the very strong increase 3 days after irradiation, which parallel senescence induction, support the hypothesis that it might be the non-necrotic,
surviving senescent cells. Most importantly, the *in vivo* relevance of our findings has to be explored. Future studies have to address the question, whether ablative irradiation can stimulate monocyte migration and intra-tumoral monocyte recruitment *in vivo*. Intriguingly, different models of sterile injury have revealed that endothelial cells and pericytes can convert the danger signals released by necrotically dying cells into cascades of chemokine gradients and adhesion molecules, which govern the recruitment of monocytes and neutrophils to the site of injury [56,57]. Along the same line, it has been shown for systemic anthracycline therapy that nucleotides released from dying cancer cells stimulate the intra-tumoral recruitment of a CD11c+CD11b+Ly6Chi monocytic cell type, which can differentiate into highly potent APCs, engulf tumor material, present it to T cells, and thereby initiate a productive anti-tumor immune response.
response [20]. Extracellular nucleotides appear to be of crucial importance in this context, since they do not only contribute to monocyte recruitment, but also support monocyte activation and differentiation as well as their intra-tumoral survival [20,58]. Hence, it is tempting to speculate that local ablative radiotherapy might be utilized for fast proliferating, hormone receptor negative, p53 mutant breast cancer in order to induce anti-tumor immune responses.

Conclusions

Here, we show that fast proliferating, hormone receptor negative breast cancer cell lines with mutant p53 intensively undergo necrosis in response to γ-irradiation, especially when applied in an ablative setting at a single dose of 20 Gy. These necrotically dying cancer cells release nucleotides, which stimulate monocyte migration and chemokinesis. In contrast, hormone receptor positive cells with wildtype p53 upregulate CD39 ectonucleotidase expression in response to irradiation, and thus do not stimulate monocyte migration. Future studies have to clarify, whether ablative radiotherapy might be utilized for local necrosis induction and concomitant nucleotide release by the dying cancer cells in order to achieve intra-tumoral monocyte recruitment, APC differentiation, and subsequent priming of adaptive anti-tumor immune responses - not only in the context of fast proliferating, hormone receptor negative and p53 mutant breast cancer.

Abbreviations

APC: Antigen presenting cell; apyrase: Nucleotide diphosphohydrolase; C12-FDG-FITC: 5-dodecanoylaminofluorescein-di[β-galactopyranoside; FCS: Fetal calf serum; FMI: Forward migration index; PI: Propidium iodide; PS: Phosphatidylserine; qRT-PCR: Quantitative real-time RT-PCR; SA-β-gal: Senescence-associated β-galactosidase; ZAV-Fmkm: Carbobenzoxy-valyl-alanyl-asparyl[O-methyl]-fluoromethylketone.

Competing interests

The authors declare no conflict of interest.

Authors’ contributions

KL, and CB conceived and designed the experiments. RH, NB, KS, and AE performed experiments and analyzed the data. KL, CB, RH, and HS wrote the manuscript. All authors discussed the results and commented on the manuscript. All authors discussed the results, read and approved the final manuscript.

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