Brain arteriovenous malformations (AVMs) are congenital abnormalities that consist of direct connections between arteries and veins, allowing high-pressure arterial blood to flow into fragile cerebral veins, resulting in a high risk of hemorrhagic stroke. The aim of AVM treatment is to prevent hemorrhage. Treatment is effective only with complete AVM removal or obliteration. Current treatments—namely surgery, endovascular occlusion, or stereotactic radiosurgery—are all associated with significant limitations. The surgical risk is generally unacceptable for lesions that are large, located in critical brain structures, or involve deep perforating arteriovenous malformations (AVMs).

OBJECTIVE Stereotactic radiosurgery (SRS) is an established intervention for brain arteriovenous malformations (AVMs). The processes of AVM vessel occlusion after SRS are poorly understood. To improve SRS efficacy, it is important to understand the cellular response of blood vessels to radiation. The molecular changes on the surface of AVM endothelial cells after irradiation may also be used for vascular targeting. This study investigates radiation-induced externalization of phosphatidylserine (PS) on endothelial cells using live-cell imaging.

METHODS An immortalized cell line generated from mouse brain endothelium, bEnd.3 cells, was cultured and irradiated at different radiation doses using a linear accelerator. PS externalization in the cells was subsequently visualized using polarity-sensitive indicator of viability and apoptosis (pSIVA)-IANBD, a polarity-sensitive probe. Live-cell imaging was used to monitor PS externalization in real time. The effects of radiation on the cell cycle of bEnd.3 cells were also examined by flow cytometry.

RESULTS Ionizing radiation effects are dose dependent. Reduction in the cell proliferation rate was observed after exposure to 5 Gy radiation, whereas higher radiation doses (15 Gy and 25 Gy) totally inhibited proliferation. In comparison with cells treated with sham radiation, the irradiated cells showed distinct pseudopodial elongation with little or no spreading of the cell body. The percentages of pSIVA-positive cells were significantly higher (p = 0.04) 24 hours after treatment in the cultures that received 25- and 15-Gy doses of radiation. This effect was sustained until the end of the experiment (3 days). Radiation at 5 Gy did not induce significant PS externalization compared with the sham-radiation controls at any time points (p > 0.15). Flow cytometric analysis data indicate that irradiation induced growth arrest of bEnd.3 cells, with cells accumulating in the G2 phase of the cell cycle.

CONCLUSIONS Ionizing radiation causes remarkable cellular changes in endothelial cells. Significant PS externalization is induced by radiation at doses of 15 Gy or higher, concomitant with a block in the cell cycle. Radiation-induced markers/targets may have high discriminating power to be harnessed in vascular targeting for AVM treatment.

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KEY WORDS phosphatidylserine externalization; endothelial cells; ionizing radiation; brain arteriovenous malformations; live-cell imaging; cell cycle; stereotactic radiosurgery; vascular disorders
ies. Endovascular treatment is usually not able to achieve a complete obliteration and is used to facilitate surgery rather than attempt cure.

Stereotactic radiosurgery (SRS), a technique to deliver focused ionizing radiation to the AVM tissues in the brain, has been widely used to treat these patients. SRS can focus on the targeted nidus precisely, with only negligible amounts of radiation delivered to the surrounding brain tissue. The margin dose of the radiation is typically 16–25 Gy, depending on the nidus size, location, shape, and proximity to critical structures. Occlusion rates are generally greater than 70%.

The purpose of SRS for AVMs is to occlude the blood vessels. The processes of AVM vessel occlusion after radiosurgery are poorly understood but are believed to involve a combination of cellular proliferation and intra-vascular thrombosis. Unfortunately, there is a delay to occlusion of 2–3 years after treatment and the risk of hemorrhage is not reduced until complete obliteration occurs. Another problem with SRS is that the radiation dose must be selected carefully so that it is not too high, which may cause radiation-induced injury to surrounding tissues, or too low, which may not be sufficient to cause obliteration of the vessels. Although SRS scoring systems have been developed and validated in clinical practice, understanding the biological response of endothelium to radiation may permit a more accurate prediction of SRS outcomes. Radiation-induced changes in some cells, such as cancer cells, have been investigated, however, the relevant studies in brain endothelial cells are limited, especially at the molecular level.

The molecular changes on the surface of AVM endothelial cells after radiation treatment may also be used as targets for vascular targeting (i.e., delivering thrombotic agents selectively to AVM sites by targeting markers on these cells, with the goal of obliterating the AVM vessels rapidly by inducing thrombosis). Although SRS scoring systems have been developed and validated in clinical practice, understanding the biological response of endothelium to radiation may permit a more accurate prediction of SRS outcomes. Radiation-induced changes in some cells, such as cancer cells, have been investigated, however, the relevant studies in brain endothelial cells are limited, especially at the molecular level.

The conventional way to study PS externalization is to use fluorescently labeled annexin V, a protein that selectively binds to PS as a reporter molecule. Normally, cultured cells or tissue slides are stained with annexin conjugates and the cells analyzed by microscopy or flow cytometry. However, because PS is a constructional lipid component of the cell membrane that is normally located on the inner surface of the membrane, many experimental treatments, such as harvest or fixation, can cause PS externalization. Hence, the results may not be reliable. Advances in microscopy have led to the development of live-cell imaging in recent years, and this represents an ideal method to observe PS externalization, because cell staining and monitoring are performed during culture. The technique allows for the visual detection of dynamic changes of PS externalization in real time after stimulation. Furthermore, it is a simple technique in which cells do not require any treatment except for the addition of fluorescently conjugated annexin V into culture medium.

Since the fluorescently labeled annexin V is added to the medium and kept there during the monitoring period without any washing steps, it is possible that background fluorescence from the dye affects the detection sensitivity. Recently, polarity-sensitive indicator of viability and apoptosis (pSIVA)-IANBD, an annexin-based, polarity-sensitive dye for PS, has become available. This indicator has no fluorescence in solution; however, it fluoresces strongly upon binding to PS on the cell surface, because of the change of its microenvironment from aqueous solution to lipid cell membrane, and this binding is reversible. In the absence of PS on the cell surface, the dye is unbound (in solution); therefore, no background fluorescence is emitted, eliminating the need to wash out the unbound dye. This makes it ideal for live-cell imaging of PS externalization. In this paper, we report our results of PS externalization of mouse brain endothelial cells after radiosurgery using the live-cell imaging with the polarity-sensitive dye pSIVA-IANBD.

Methods

Materials

Black, 96-well cell-culture plates with a flat and clear bottom were obtained from Ibidi GmbH. The bEnd.3 cells were from the American Type Culture Collection, and pSIVA-IANBD apoptosis/viability microscopy kits were purchased from Imgenex. Attachment factor for cell culture was a product of Cell Systems. DMEM, fetal bovine serum (FBS), and 1% penicillin/streptomycin were from Invitrogen.

Cell Culture

bEnd.3 cells, an immortalized cell line generated from mouse brain endothelium, were grown according to the supplier’s instructions in high-glucose DMEM containing 10% FBS, 4.5 g/L glucose, 3.7 g/L sodium bicarbonate, 4 mM glutamine, 100 U/ml penicillin, and 100 μg/ml streptomycin, at a pH of 7.4. Cells were maintained in a humid chamber at 37°C in an atmosphere of 95% air and 5% carbon dioxide in 75-cm² tissue-culture flasks. Confluent flasks were trypsinized and around 0.2% of the cells in
300 μl culture medium were seeded into individual wells (with a culture area of 0.55 cm²) of a 96-well cell-culture plate precoated with attachment factor. The cells were allowed to grow for 1 day (about 20%–30% confluence) before undergoing irradiation.

**Cell Irradiation**

A linear accelerator (LINAC; Elekta AB) at Macquarie University Hospital was used to irradiate the cells. The 96-well plate was CT scanned and a radiation-dose distribution was planned on the Elekta XiO planning system to deliver a differential dose (25 Gy, 15 Gy, 5 Gy, and < 0.5 Gy) to each of the 4 quadrants of the 96-well plate. Irradiation of the cells was carried out using 6-MV photons from anterior- and posterior-directed radiation fields. The cells in the area that received < 0.5 Gy were treated as a sham-radiation control. The culture medium was changed just before irradiation.

**Live-Cell Imaging**

After irradiation, the dyes pSIVA-IANBD (1.5 μl) and propidium iodide (PI, 0.75 μl), in the apoptosis/viability microscopy kit, and the apoptosis-inducing agent camptothecin (final concentration 10 μM) were added to the corresponding wells. The 96-well plate containing the cells was then transferred to the automated stage of a Nikon Ti inverted epi-fluorescent microscope (Nikon Corp.) equipped with an environmental chamber set at 37°C. Humidified 5% carbon dioxide was supplied to the wells for the duration of the experiment. A x40 objective (NA0.6) was used to collect transmission images, while pSIVA was detected in the FITC fluorescent channel (Ex 485 Em 521/13) and PI was detected in the Texas Red channel (Ex 607 Em 607/18). Two fields of view were selected in each well of triplicate wells, generating 6 fields of view for each radiation-dose group. Images were taken every 30 minutes at each field of view for 72 hours. The first image was taken 4 hours after irradiation. The experiment was repeated twice.

**Detection of PS Externalization and Cell Death**

Manual counting by 2 of the authors independently (1 of whom was blinded to sample identity) of cell number and pSIVA- and PI-positive staining spots was performed in each of the 6 fields of view in each group to calculate the degrees of PS externalization and cell death for the cells exposed to different doses of radiation. Counting was performed on images taken at 4, 24, 49, and 74 hours after irradiation. The results shown are a representative analysis and are expressed as the ratios of positive counting/total cell number.

**Flow Cytometric Analysis of Cell Cycle**

bEnd.3 cells were seeded into 75-cm² culture flasks, allowed to grow for 1 day (about 30% confluence), then irradiated using the linear accelerator, whereby cells received either 5-, 15-, or 25-Gy dose of radiation. Control cells were not exposed to radiation. Cells were harvested 3 days later, fixed in ice-cold 70% ethanol, and stained with PI (10 μl PI in 1 ml PBS containing 0.05% Triton X-100 and 100 μg ribonuclease A) for 30 minutes for detection of DNA. Flow cytometry analysis of cell cycle was performed on an LSR Fortessa X-20 cytometer (Becton, Dickinson and Co.). Data were acquired with DIVA software (version 8.1, Becton, Dickinson and Co.), collecting 10,000 events per sample with PI area fluorescence detected on a linear scale, and % forward (FWD)-area set to 0.40. Data were analyzed with CellQuest Pro (version 6.0; Becton, Dickinson and Co.), and with Modfit LT (version 3.2) for cell cycle analysis gating on single cells defined as those with proportional FWD-height versus FWD-area. The composite data figure was prepared using Adobe Illustrator CS6 (version 16.0.0; Adobe Systems Inc.).

**Statistical Analysis**

Results are expressed as mean ± SD. Comparison of differences between test groups was made by unpaired two-tailed Student t-test. Differences were considered significant at p < 0.05.

**Results**

**Effect of Radiation on Cell Proliferation**

The cell confluence at the time of radiation treatment was 20%–30%, leaving enough room for the cells to proliferate during the 3-day incubation period under the microscope. The sham-radiated cells grew well, demonstrating the conditions in the chamber were appropriate for the cells. The cell number in each field of view was between 8 and 30.

The cell proliferation rates in the wells were radiation-dose dependent. The results were similar in the 2 separate experiments, and the rates from a representative run are shown in Fig. 1. In the sham-radiation wells, the number of cells in the fields of view increased by 140% by Day 3. Using this result as a control, radiation at a dose of 5 Gy significantly reduced the cell proliferation rate. The increase in cell number at Day 3 was less than 50% (p < 0.01). The higher radiation doses (15 Gy and 25 Gy) totally inhibited cell proliferation. The cell number was eventually reduced by around 20% at Day 3. The differences in percentage changes of cell numbers at Day 3 were not statistically significant between the groups receiving 15- or 25-Gy doses (p = 0.72), but both groups were significantly lower than the group treated with the 5-Gy dose and the sham-radiation controls (p < 0.01). The percentage change of cell number at Day 3 in the camptothecin-positive control group was similar between the groups treated with 15- or 25-Gy doses but significantly lower than the group treated with the 5-Gy dose and the sham-control groups (p < 0.05).

**Morphology Changes of the Cells After Irradiation**

The low-density bEnd.3 cells at the time of irradiation exhibited a normal endothelial cell morphology with flattened and epithelial appearance, and some were elongated like fibroblasts (Fig. 2A and C). In comparison with sham radiation–treated cells, the irradiated cells showed distinct pseudopodial elongation with little or no spreading of the cell body and more cells with increased cytoplasmic volume. This was particularly apparent at late stages of the experiment and with higher radiation doses (Fig. 2C).
PS Externalization and Cell Death After Irradiation

In cells that underwent PS staining using pSIVA, only a few green spots could be seen per field of view at the beginning of imaging (4 hours after radiation exposure), and no significant differences in PS externalization could be detected in irradiated cells at any radiation-dose level in comparison with sham-radiation control cells (Fig. 2A and B). However, the percentages of pSIVA-positive cells, represented by the ratio of green spots to cell number in the corresponding fields, were significantly higher (p = 0.04) in the cells that received a 25- or 15-Gy dose of radiation 24 hours after treatment (Figs. 2A, 2B, and 3A). This effect was sustained until the end of the experiment (3 days), with the highest ratios at the last time point (p < 0.01). Radiation at 5 Gy did not induce significantly higher PS externalization compared with the sham-radiation control at any time points (p > 0.15). The effect of camptothecin on cell PS externalization was similar to that seen with the 25- and 15-Gy radiation doses. However, in comparison with radiation-treated cells, the increase in the pSIVA-positive cells in the second half of the incubation period was more notable in the camptothecin-treated cells. The results from a representative run are shown in Figs. 2 and 3.

The percentages of PI-positive cells, represented by the ratio of red spots to cell number in the corresponding fields, had a similar trend with the pSIVA-positive cells, although the values were much lower in all the radiation-dose groups at all time points (Figs. 2A and 3B). The percentages were significantly higher in 25- and 15-Gy treated cells at 24 hours after irradiation and later time points. In the cells treated with a 5-Gy dose of radiation and camptothecin, the percentage of PI-positive cells was significantly higher than sham-radiation control cells only at the end of the experiment (Figs. 2A and 3B).

Statistical analysis did not show any significant differences between 25- and 15-Gy radiation doses in cell pSIVA staining and PI staining at any time point. However, there were significant differences in pSIVA and PI staining between 15- and 5-Gy radiation doses at 49 and 74 hours postirradiation.

Cell Cycle Analysis of the Irradiated Endothelial Cells

Irradiated bEnd.3 endothelial cells were also analyzed by flow cytometry. Here, the effect of irradiation was clearly evident, as even low-dose radiation (e.g., 5 Gy) resulted in a change in the cells FWD-area versus FWD-height dot-plot profile (Fig. 4A). Moreover, even in a simple ungated analysis, 15- and 25-Gy radiation doses appeared to alter the proportion of cells in G1 versus G2 of the cell cycle (Fig. 4A). These doses of radiation also more than doubled the number of polyploid cells by 3 days (events right of the 2N peak in Fig. 4A). To confirm these results, a proper cell cycle analysis was performed by first gating on single cells, defined as cells that had a proportional FWD-height versus FWD-area ratio (region 1; R1, blue gate). The cell cycle analysis indicated that 48% of the cells had accumulated in the G2 phase of the cell cycle after a 25-Gy radiation dose, compared with just 19% in nonirradiated cells, with a similar effect after a 15-Gy radiation dose (23% G2) (Fig. 4B). Taken together, these data indicate that irradiation induces the growth arrest of bEnd.3 cells, with cells accumulating in the G2 phase of the cell cycle and with many cells becoming senescent as evident by their polyploid state.
The effect of ionizing radiation on cells is well documented. It includes 2 main aspects: direct cellular damage through modification of macromolecules such as DNA and proteins, and indirect cellular damage through production of reactive oxidative species. Depending on the level of damage, cell cycle arrest, DNA repair, senescence, or apoptosis will be initiated. Endothelial cells are very sensitive to radiation. Our previous studies revealed proinflammatory and thrombotic molecule changes on the surface of brain endothelial cells after ionizing radiation exposure. The current study further confirmed that radiation at clinically relevant doses (15 and 25 Gy) has a clear effect on the bEnd.3 cells. Radiation at both doses totally inhibited the growth of the cultured cells, caused growth arrest and remarkable morphological changes, and induced significant PS externalization. Although the lower radiation dose (5 Gy) did not cause clear morphological changes or induce significant PS externalization, it clearly inhibited the growth of these cells.

Live-cell imaging takes advantage of the ability to sample data from essentially the same cells (identical fields of view) at different points in time postradiation exposure. This allowed for easy tracking of proliferation of individual cells but only for a limited number of cells (i.e., those within the imaged fields). However, flow cytometry analysis permits analysis of significantly more cells (10,000 cells or more) and additionally defined that irradiation caused the accumulation of cells in the G2 phase of the cell cycle. The results are consistent with previous reports on ionizing radiation effect on cells. Thus, the radiation induced a growth inhibition that was largely due to cell cycle inhibition, senescence, and/or cell death.

Since the distribution of PS on the cell surface is uneven and apoptotic cells may break into several pieces,
May be adequate to induce the necessary changes in externalization. The results suggest that the dose of 15 Gy cells in cell proliferation, morphological changes, and PS effects of 15- and 25-Gy radiation doses on the bEnd.3 endothelium of A VM tissues can be achieved in vivo (data not shown). Preliminary data suggest that selective radiation-induced marker for vascular targeting, although more work is required to find the optimal radiation dose for vascular targeting. Given that high doses of radiation, including current clinical doses, can lead to off-target effects and late radiation necrosis, it would be clinically very significant if lower radiation doses could induce sufficient molecular changes on the endothelial surface for vascular targeting.

Live-cell imaging can monitor molecular changes on the cell surface in real time without the need to process the cells except for the addition of the dyes into the culture medium. It is designed to observe the same sets of cells in each field of view at different time points, eliminating the experimental errors caused by the variation among the cells. It also has the capacity to monitor the changes of a single cell. Up to a few hundred fields of view across the cell culture plate can be monitored in just 1 experiment. With the radiation dose designed to deliver the desired doses to different parts or wells of the plate, this technique can detect the effects of different radiation doses on the cells in real time with just 1 run. However, there were some drawbacks with the system that we used.

The bright-field image quality was affected near the edges of the narrow wells, and condensation on the lid altered the brightness of the images over time. In this system, the environmental chamber lid could not be removed once mounted onto the microscope; therefore the culture medium could not be changed during the imaging process, limiting the monitoring period to a maximum of 3 days in this study. The results cannot reveal whether all the PS-positive cells will eventually die or can recover or be rescued to normal cells. The expression of PS on the cell surface increased with time during the monitoring period. No peak expression time was identified. Nevertheless, large amounts of information regarding the changes of the cells during the monitoring period can be obtained if the conditions are carefully controlled and appropriate control cells are included in the experiments.

Naturally, it is a limitation of all in vitro investigations that the microenvironment for the cultured cells is different from the in vivo conditions. There are no other cells to interact with these cells, such as macrophages to get rid of abnormal cells. The cells in vivo do not have the space to proliferate. Nevertheless, this in vitro study provides valuable insight into what might happen in vivo and, thus, forms a strong foundation for designing in vivo work in the future.

Conclusions

Ionizing radiation causes remarkable cellular changes in endothelial cells. Significant PS externalization is induced by radiation at doses of 15 Gy or higher, concomitant with a block in the cell cycle. Radiation-induced markers/targets may have high discriminating power to be harnessed in vascular targeting for AVM treatment.
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Fig. 4. Flow cytometry analysis of irradiated bEnd.3 cells. A: The proportional size (FWD-H [forward height] and FWD-A [forward area]) and nuclear content represented by intracellular PI fluorescence of permeabilized bEnd.3 cells 3 days after irradiation. B: Cell cycle analysis of single cells (R1 gate, as defined in A) after irradiation, indicating the percentage of cells in the G1, S, or G2 phase of the cell cycle. Data shown are representative of independently repeated experiments.

Disclosure
The authors report no conflict of interest concerning the materials or methods used in this study or the findings specified in this paper.

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Conception and design: Zhao, Johnson, Grace, Ukath, Stoodley. Acquisition of data: Zhao, Johnson, Chen, Grace, Ukath, Lee, McRobb, Sedger. Analysis and interpretation of data: Zhao, Johnson, McRobb, Sedger, Stoodley. Drafting the article: Zhao, Sedger. Critically revising the article: Johnson, Lee, McRobb, Sedger, Stoodley. Reviewed submitted version of manuscript: all authors. Approved the final version of the manuscript on behalf of all authors: Zhao. Statistical analysis: Zhao. Study supervision: Zhao, Stoodley.

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