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A Zebrafish Live Imaging Model Reveals Differential Responses of Microglia Toward Glioblastoma Cells In Vivo

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Abstract

Glioblastoma multiforme is the most common and deadliest form of brain cancer. Glioblastomas are infiltrated by a high number of microglia, which promote tumor growth and surrounding tissue invasion. However, it is unclear how microglia and glioma cells physically interact and if there are differences, depending on glioma cell type. Hence, we have developed a novel live imaging assay to study microglia–glioma interactions in vivo in the zebrafish brain. We transplanted well-established human glioblastoma cell lines, U87 and U251, into transgenic zebrafish lines with labelled macrophages/microglia. Our confocal live imaging results show distinct interactions between microglia and U87, as well as U251 glioblastoma cells that differ in number and nature. Importantly these interactions do not appear to be antitumoral as zebrafish microglia do not engulf and phagocytose the human glioblastoma cells. Finally, xenotransplants into the \textit{irf8}⁻/⁻ zebrafish mutant that lacks microglia, as well as pharmacological inhibition of the CSF-1 receptor (CSF-1R) on microglia, confirm a prominent role for zebrafish microglia in promoting human glioblastoma cell growth. This new model will be an important tool for drug screening and the development of future immunotherapeutics targeting microglia within glioma.

Keywords: microglia, glioma, brain tumor, macrophages

Introduction

Gliomas are the most common type of malignant brain tumor. The most aggressive and frequently occurring form of glioma is known as glioblastoma multiforme (GBM), or high grade astrocytoma. GBM is incurable and has a median survival rate of less than 14 months.1,2 GBMs display extensive intra- and intertumoral heterogeneity in both morphological features and genetics.3,4 Furthermore, glioblastomas show a diffuse appearance, which is the main reason why these tumors cannot be cured by surgical resection. Although Scherer described the migration of glioma cells away from the main tumor mass as early as 1940,5 the underlying mechanisms are still not understood. Recent studies suggest that high levels of macrophages/microglia within the glioma are positively correlated with glioma grade and invasiveness.6–17

Microglia are the residential macrophages of the brain.18,19 Among many other functions, microglia sense injuries in the brain immediately, migrate to the site of the lesion, and phagocytose detrimental material.20–23 Strikingly, a comparable course of events is observed during glioma progression in the brain, where microglia and infiltrating macrophages are attracted to and colonize the tumor.6,7,9,16,17 However, instead of antitumoral activity they display protumoral functions and promote tumor growth.11–15,24–27 Microglia/macrophages within glioblastomas have been found to account for 30% of the glioma mass and have been implicated in establishing an immunosuppressive microenvironment.6,7,16

Thus, understanding the nature of microglia–glioma interactions in vivo could be the first step to develop future alternative strategies to interfere with glioma growth and invasiveness.

The zebrafish represents a powerful model system to explore cellular responses and molecular events in vivo. It has been established as a model to study various types of human cancer, ranging from B-Cell/T-Cell leukemia and melanoma to glioma.28–38 We have utilized a zebrafish xenotransplantation live imaging model to address microglia–glioma interactions. The zebrafish larva provides optimal characteristics that are advantageous for these studies. First, the zebrafish immune system is unique in the sense that after fertilization, the larvae survive only with the innate immune system.39–40 Maturation of the immune system leading to the development of the adaptive immune response occurs at
between 3 and 6 weeks postfertilization.\textsuperscript{39,40} Thus upon xeno-
graft these early events during tumor colonization can be
studied in detail without interference by the highly diver-
sified and complex response of an adaptive immune system.
Second, a major benefit of the larval model is the optical
transparency, which makes it possible to directly observe and
classify the different microglia–glioma interactions \textit{in vivo} in
high resolution. To perform similar studies in a rodent model
the insertion of a cranial window is necessary.\textsuperscript{41} While fea-
sible, this requires an additional surgical procedure that needs
to be tolerated by the animal. Furthermore, immunosup-
pression has to be applied upon transplantation of human
cells, which might impact microglia–glioma interactions as
well. To overcome this limitation, orthotopic syngeneic
mouse models like the GL261 glioma model have been
developed.\textsuperscript{42} This model, in combination with two-photon
imaging, has been used very recently to monitor how mi-
croglia respond to mouse GL261 glioma cells.\textsuperscript{43–45} However,
interactions of microglia with human glioblastoma cells have
never been visualized to date.

We have exploited recently established zebrafish lines with
fluorescently labelled macrophages/microglia to simultaneously
monitor the migration and movement of microglia and gli-
blastoma cells, as well as their interactions with each other.
Transplantation of human U87 and U251 glioblastoma cells
into the zebrafish brain led to an immediate microglial re-
sponse. To test if these responses were specific for glioblas-
toma cells, we performed heterotopic transplants of human
fibrosarcoma cells (HT1080). Interestingly, we observed spec-
cific nonphagocytic interactions with U87 and U251 cells,
which were different in number and in nature. Importantly,
microglial responses toward HT1080 cells were very different
and many of these cells were immediately engulfed. Finally,
xenotransplants into the \textit{irf8}\textsuperscript{−/−} zebrafish mutant, which lacks
microglia, confirmed a prominent role for microglia in pro-
moting U87 and U251 tumor cell survival. In summary, our
results show that the zebrafish larva is a powerful tool to study
specific interactions between microglia and glioma cells.

**Materials and Methods**

**Cell culture**

Human U87MG glioblastoma cells were kindly provided by
Prof Tobias Pukrop (University Hospital Regensburg,
Germany). Human U251MG glioblastoma cells were pur-
chased from CLS Cell Lines Service (Eppelheim, Germany),
and human HT1080 cells were kindly provided by Dr Pamela
Brown (SURF, University of Edinburgh). U87MG cells were
cultured in Dulbecco’s modified Eagle’s medium (DMEM)
containing 1% \textit{l}-glutamine and supplemented with 1% (v/v)
Penicillin/Streptomycin (100 mg/mL penicillin and 100 mg/
l streptomycin) and 10% (v/v) fetal calf serum at standard
conditions of 100% humidity and 5% CO\textsubscript{2}, at 37°C. U251MG
cells were cultured in DMEM containing 1% \textit{l}-glutamine and
supplemented with 1% (v/v) Penicillin/Streptomycin, 0.1 mM
nonessential amino acids, 1 mM sodium pyruvate, and 10% (v/v)
fetal calf serum at standard conditions of 100% humidity and 5%
CO\textsubscript{2}, at 37°C. HT1080 cells were cultured in DMEM contain-
ing 30 mg/L GlutaMAX, 4.5 g/L D-Glucose, and 2.5 mM HEPES
supplemented with 1% (v/v) Penicillin/Streptomycin and 10%
(v/v) fetal calf serum at standard conditions of 100% humid-
ity and 5% CO\textsubscript{2}, at 37°C. Cells were harvested on the day of
the xenograft transplantation. The cells were washed with
phosphate-buffered saline (PBS) and detached from the surface
of the flask by incubation in 2 mM EDTA in PBS for 15 min at
34°C. The EDTA was removed with a 10 mL DMEM wash
followed by a 10 mL PBS wash; cells were centrifuged for
1.5 min at 0.2 g between washes. The resulting cell pellet was
resuspended in 100 μL DMEM, supplemented with phenol red
(1:10 v/v), to a final concentration of 5 × 10\textsuperscript{3} cells/mL in prep-
paration for the xenografts.

**Lentiviral transductions and CM-Dil dye labelling**

For lentiviral transductions, cells (4.5 × 10\textsuperscript{5}) were seeded
into 60 mm dishes in a final volume of 5 mL DMEM. Cells
were allowed to adhere to the dish for 4 h, at standard incubator
conditions. HT1080 cells were grown on Matrigel to allow
transformation. Polybrene (5 ng/mL) was added directly to the
culture medium and incubated for 10 min at 37°C. The Lv-
cpt-IRES-mCherry-opre viral vector (Dr. P. Brown, SURF)
at a multiplicity of infection of 10 was then added to the ex-
isting media and was incubated at standard conditions for 48 h.
After 48 h, the media containing viral vector were removed.
The transformed cells were detached from the culture dish by
15 min of 2 mM EDTA-PBS treatment at 37°C and given a
10 mL DMEM wash and then 10 mL PBS wash; cells were
centrifuged for 1.5 min at 0.2 g between washes. The resulting
cell pellet was resuspended in 10 mL DMEM and seeded into
cell culture dishes at the required density. This procedure
gave rise to cell lines that stably expressed the mCherry reporter
protein, which was evenly distributed throughout the cells.
These mCherry expressing cells are referred to as U87, U251,
and HT1080 cells throughout the text.

CM-Dil Dye (Molecular Probes) was dissolved in DMSO
according to manufacturer’s instructions, and a working
concentration of 2 μM was used for cell labeling. Cells were
incubated in 2 μM CM-Dil Dye for 10 min at 4°C. This was
followed by two PBS washes and a media replacement in
preparation for harvesting.

**Mammalian cell growth kinetics**

The growth rates of HT1080 mCherry, U87 mCherry, and
U251 mCherry cells at 34°C compared to 37°C were deter-
mined over a 4-day period. Monolayer cultures of each cell
line were harvested, using Accutase (Sigma-Aldrich), when
~ 75% confluent. The detached cells were washed once with
1 × PBS and resuspended in culture medium. The cells were
counted using a Neubauer counting chamber (depth 0.1 mm,
1/400 mm\textsuperscript{2}; Hawksley), and 1500 cells, in 1 mL, of culture
medium were transferred into the wells of 12-well plates.
Cells were cultured continuously for up to 4 days, without
replenishing media, at either 34°C or 37°C under normoxic
conditions. Each day (at 24-h intervals), the cell number, in
three individual wells, was counted for each cell line and the
average cell number was calculated. The experiment was
repeated in triplicate, and the average cell numbers, from the
three experiments, were plotted to produce growth curves.

**Zebrafish maintenance, xenografts, and pharmacological treatments**

Zebrafish were housed in a purpose built zebrafish facility,
in the Queen’s Medical Research Institute, maintained by the
University of Edinburgh Biological Resources. All zebrafish
larvae were kept at 28°C on a 14-h light/10-h dark photoperiod. Embryos were obtained by natural spawning from adult mpeg1:EGFP<sup>46</sup> and irf8<sup>−/−</sup> zebrafish strains. Animal experimentation was approved by the Ethics Review Committee of the University of Edinburgh and the Home Office, in accordance with the Scientific Procedure Act 1986. Embryos were treated with 200 μM 1-phenyl 2-thiourea (PTU) from the end of the first day of development for the duration of the experiment to inhibit pigmentation. Xenografts were obtained from the end of the first day of development for the duration of the experiment to inhibit pigmentation. Xenografts were conducted on larvae that had been dechorionated and anesthetized with 2.5 mM Ethyl 3-aminobenzoate methane sulfonate (Tricaine) in zebrafish embryonic medium (6.4 mM KCl, 0.22 mM NaCl, 0.33 mM CaCl<sub>2</sub> 2H<sub>2</sub>O, 0.33 mM MgSO<sub>4</sub> 7H<sub>2</sub>O), at 3 dpf. Anesthetized zebrafish larvae were immobilized in 1.5% (w/v) low-melting point agarose solution made with zebrafish embryonic medium. The heads of the embryos were carefully exposed from the agarose before transplantation (Fig. 1). All transplantations were performed using a Femtojet 4i microinjector (Eppendorf, Hamburg, Germany). Transplantation needles were prepared from borosilicate glass capillaries, lacking an internal filament, using a glass micropipette puller (Sutter Instrument Co. P-87, Novato). U87/251/HT1080 cells (n = 8–30) were transplanted into the optic tectum of each zebrafish larva. Individual zebrafish larvae containing U87/251/HT1080 cells were maintained in 12-well plates, containing 4 mL zebrafish embryo medium supplemented with 200 μM PTU, at 34°C. For pharmacological treatments, U87 cell transplanted larvae were maintained in 24-well plates in zebrafish embryo medium supplemented with 200 μM PTU, at 34°C. Transplanted larvae were treated from 4 dpf (1 day posttransplantation) with either 1% DMSO (control) or 1% DMSO/25 μM BLZ945 for 5 days. Larvae were imaged at 4 dpf before the start of treatment to obtain microglial cell counts and U87 cell counts. Treated larvae were kept individually and imaged at the end of treatment (9 dpf) to obtain final microglia and U87 cell counts.

Imaging

Zebrafish larvae were anesthetized with 2.5 mM tricaine in embryonic medium and mounted in 1.5% (w/v) low-melting point agarose. Time lapse and still images were obtained using an Andor spinning disk confocal fluorescent microscope with a 20 × /NA 0.75 Olympus objective lens. The acquired z stacks for both single time point and time lapse images were selected to encompass all regions of interest within the zebrafish larval brain. Time lapse imaging was conducted for a period of 14–17 h with the time interval between images set at 3 min. Single time point images were obtained at 0–4 days posttransplantation (dpt).

Analysis

Analysis of all images was conducted using Imaris (Bitplane, Belfast, United Kingdom). All data were plotted using GraphPad and analyzed using one way ANOVA and post hoc analysis with Sidak’s multiple comparisons test. The cell number was determined for each image by manual counting in 3D. These values were used to plot survival curves for each cell type in the different zebrafish lines. The survival curve was determined as the ratio expressed as a percentage between the cell number at each time point and the starting cell number.

The number of microglia interactions between individual microglia and the transplanted cells was measured by counting the number of direct interactions between both cell types and normalizing for the number of xenografted cells at the respective time point. Interactions were defined as direct contacts between microglial surfaces and transplanted cell surfaces, including engulfment and phagocytosis of transplanted cells (Supplementary Fig. S1B–E; Supplementary Data are available online at www.liebertpub.com/zeb). If a microglial cell showed several protrusions in contact with a single transplanted cell, this was counted as one interaction between these two cells.

The invasiveness of the different cell types was determined by measuring the spread of fluorescently labelled cells in 3D within the zebrafish larval brain over time. Using Imaris, a 3D sphere was created to encompass all fluorescent cells of interest. Transplanted cells that had been engulfed by macrophages/microglia were not included within the sphere (see Results section; Supplementary Fig. S2C, D). The diameter of the created 3D sphere was used to measure invasiveness. Invasiveness was determined as a percentage between the...
final diameter of the sphere, required to encompass the transplanted cells 4 dpt compared to the diameter of the sphere surrounding the initial cell mass directly after transplantation.

Results

In vivo imaging reveals different growth patterns and microglial responses for U87 and U251 glioblastoma cells in the larval zebrafish brain

The combination of fluorescent transgenic tools and the innate optical transparency of the zebrafish larva makes it possible to study microglial interactions under high temporal and optical resolution in the living brain. To study microglial responses to xenografted human glioblastoma cells, we made use of the mpeg1:EGFP transgenic zebrafish, in which all macrophages, including microglia, can be visualized and tracked. The xenotransplantation protocol used in this study is an optimized version of our previously developed injection protocol and is minimally invasive. Briefly, larvae were mounted in low-melting point agarose, and their heads were carefully exposed from the agarose using a sharp needle before transplantation (Fig. 1A–A2). Transplantations were performed into the left optic tectum using a glass capillary (Fig. 1B–B3). Sham injections of culture media led only to a short injury response of microglial cells that were in direct proximity to the injection site. This injury response was usually resolved within a few hours (data not shown). To study microglial responses to glioblastoma cells, we transplanted human-derived U87mCherry (referred to as U87) and U251mCherry (referred to as U251) glioblastoma cells into the optic tectum and whole brain confocal imaging was performed at 0–4 dpt. To test if microglial responses were specific for glioblastoma cells, we performed heterotopic transplants of the human fibrosarcoma cell line HT1080mCherry (referred to as HT1080) and performed imaging at the same time points.

Upon transplantation of U87 and U251 glioblastoma cells into the larval zebrafish tectum, the cells invaded the surrounding tissue of the optic tectum moving away from the initial transplantation site (Fig. 2). However, it was immediately obvious that U251 cells and U87 cells had very different cellular morphologies within the zebrafish brain (Fig. 2). U251 glioblastoma cells showed infiltrative cellular protrusions that projected from the main cell bodies into the surrounding tissue (Fig. 2B/C, arrows). In most U251 samples, it was observed that these projections extended to over 100 μm in length and formed interactions with microglia outside of the central glioma mass (Fig. 2C, arrows, arrowheads). Furthermore, U251 cells attracted many microglia that stayed within the vicinity of the U251 cells over the duration of the experiment (Fig. 2A–C). These microglia showed different types of interactions with U251 cells (see below) that could be observed over the full duration of the experiment (Fig. 2A–C, arrowheads).

Transplanted U87 cells showed a different morphology compared to U251 cells (Fig. 2D–F). The borders of the main U87 glioblastoma cell mass were more defined, and cellular protrusions were observed to a much lesser degree compared to the U251 cells (compare Fig. 2C, arrows to Fig. 2F, arrows). Nevertheless, single-invasive U87 cells were observed outside the main cell mass (Fig. 2E, F). U87 cells also attracted microglia and interactions between the U87 cells and microglia (see below) were observed at all time points (Fig. 2D–F, white arrow heads).

Interestingly, heterotopic transplants of human HT1080 cells into the larval zebrafish brain resulted in a different outcome. HT1080 cells attracted many microglia (Fig. 2G) and these microglia immediately engulfed HT1080 cells (Fig. 2H, I and Supplementary Fig. S1B). Thus after 4 dpt, many HT1080 cells were detected within phagosomes of microglia, while less than half of the HT1080 cells escaped engulfment by microglia (Fig. 2I and Supplementary Fig. S1B).

Microglia differently interact with U87 and U251 cells

To analyze the nature of interactions between microglia and the different transplanted cell types in more detail, we performed live imaging of U87, U251, and HT1080 transplanted mpeg1:EGFP fish at 3-min intervals for up to 14 h. These time series revealed obvious differences in the responses of microglia toward the different cell types. Interestingly, microglia behavior was very different upon transplantation of U87 and U251 cells. U251 cells immediately attracted many microglia (Fig. 3A and Supplementary Movie S1). These microglia deeply infiltrated the U251 cell mass and exhibited continuous interactions over the remaining hours (Fig. 3A and Supplementary Movie S1). We observed two main types of cellular surface interactions. The first type of interaction observed was a long-lasting interaction between individual microglia and single U251 cells (Fig. 3A–A10, [asterisk, arrows] and Supplementary Movie S1). The second type of interaction observed was of dynamic nature, reflected in constant extensions and retractions of microglial processes that made contact with the surface of U251 cells (Fig. 3A–A10, [asterisk, arrows] and Supplementary Movie S1). These contacts were of short duration (minutes) and individual microglia interacted with several U251 cells. In comparison to U251 cells, U87 cells attracted a lower number of microglial cells and they did not penetrate the U87 cellular mass (Fig. 3B and Supplementary Movie S2). Despite this difference, we detected interactions between the microglia and transplanted U87 cells. Microglia remained at the margin of the U87 cell mass and sent out and retracted cellular protrusions to make contact with the U87 cells (Fig. 3B–B5, [asterisks, arrows]). As observed for U251 cells, these contacts were of short duration and individual microglia interacted with several U87 cells (Supplementary Movie S2).

As stated before, the microglial response toward HT1080 cells was very different. Following transplantation of HT1080 cells into the larval zebrafish brain, the main microglial response resulted in the immediate engulfment of many of these cells (Fig. 3C and Supplementary Movie S3). Over the time course, several microglia cells were found to contain HT1080 cellular fragments within their phagosomes (Fig. 3C–C6, [arrowheads] and Supplementary Fig. S1B–SB2). This explains the decrease in HT1080 cell numbers that we observed over the 4-day time course (Fig. 4A).

U87 and U251 glioma cells show differences in growth and number of microglia interactions in the larval zebrafish brain

To investigate the growth of the different cell types upon transplantation, we performed cell counts based on the confocal
FIG. 2. U251, U87, and HT1080 cells exhibit different growth patterns in the zebrafish optic tectum and stimulate an intensive microglia response. Representative confocal images of the optic tectum of mpeg1:EGFP zebrafish transplanted with U251mCherry (U251) glioblastoma cells, U87mCherry (U87) glioblastoma cells, and HT1080mCherry (HT1080) fibrosarcoma cells at 3 dpf. (A–C) Images from top to bottom are in chronological order showing 0, 2, and 4 dpt of U251 cells; (A’–C’) Higher magnification of regions of interest in (A–C) to highlight interactions of microglia (green) and U251 cells (red). U251 cellular projections are marked with a white arrow. Microglia interacting with U251 cells are marked with a white arrowhead. (D–F) Images from top to bottom are in chronological order showing 0, 2, and 4 dpt of U87 cells; (D’–F’) Higher magnification of regions of interest in (D–F) to highlight interactions of microglia (green) and U87 cells (red). U87 cellular projections are marked with a white arrow. Microglia interacting with U87 cells are marked with a white arrowhead. (G–I) Images from top to bottom are in chronological order showing 0, 2, and 4 dpt of HT1080 cells; (G’–I’) Higher magnification of regions of interest (G–I) to highlight interactions of microglia (green) and HT1080 cells (red). Microglia interacting with HT1080 cells are marked with a white arrowhead. Scale bars for (A–I): 50 µm. Scale bars for (A’–I’): 30 µm. All images represent maximum intensity projections of confocal stacks. Images were captured using an Andor spinning disk confocal microscope with a 20×/NA 0.75 objective. Insets in (A, G, and D) show the orientation of samples (anterior to the top) and the region imaged (white rectangle). dpf, days postfertilization; dpt, days posttransplantation.
z-stack images acquired at the different time points. These results show that U87 survival was significantly higher at all time points compared to both U251 and HT1080 (Fig. 4A). Over the course of the experiment, U87 cell numbers increased to 180% - 35.1%, while U251 cell numbers remained constant (Fig. 4A). The observed difference in growth for U87 cells and U251 cells upon transplantation is in line with their in vitro growth rates. The comparison of growth rates for these cells at 37°C and 34°C in vitro showed that U87 cells grow much faster than U251 cells at both 37°C and 34°C (Supplementary Fig. S1A). In comparison, HT1080 cell numbers decreased to 47.6% - 4.8% at 4 dpt in the zebrafish larval tectum (Fig. 4A). This is in contrast to their
growth in vitro as HT1080 cells were observed to grow faster than U251 cells at 37°C and 34°C (Supplementary Fig. S1A). Thus, orthotopic transplants of U87 and U251 glioblastoma cells into the optic tectum of the zebrafish brain show a greater rate of survival compared to heterotopic HT1080 transplants.

Although glioma infiltration by microglia has been described in many studies, interactions between microglial cells and the glioma cells have only been described recently in vivo for a mouse GL261 glioma model. Microglial interactions with human glioblastoma cells have never been directly visualized in vivo at high resolution until now. To ascertain if the different transplanted human glioma cell types stimulate a different number of microglial interactions, the number of individual microglia cells in direct contact with transplanted cells was analyzed based on the confocal stacks acquired between 0 and 4 dpt (see Materials and Methods section and Fig. S1C, E).

U251 glioblastoma cells stimulated a significantly higher number of microglial interactions compared to U87 (Fig. 4B, C). One way ANOVA of the total number of microglial interactions over the whole 4-day period indicated a significant difference in the number of microglial interactions with U251 cells (2.61 ± 0.150, n = 15) compared to U87 cells (1.73 ± 0.264, p = 0.037, n = 8) (Fig. 4C). HT1080 cells stimulated a high number of microglial interactions (3.31 ± 0.338, n = 10; Fig. 4B, C) compared to U87 (1.73 ± 0.264, p = 0.0005, n = 8). However, microglia-HT1080 cell interactions were very different in nature compared to interactions with U87 and U251 cells and included a high degree of engulfment of HT1080 cells.

We have shown that transplanted U87, U251, and HT1080 cells differ in survival rates and microglial responses. To test if these cell types show differences in their infiltrative behavior, we measured their invasiveness in 3D over time. This was achieved by creating a 3D sphere that accurately encompassed the transplanted cells at the different time points (see Materials and Methods section and Supplementary Fig. S2D). The diameter of the created 3D sphere was used to calculate cell invasiveness in percent. All error bars represent the standard error of the mean.
to measure invasiveness. By calculating the fold increase between spheres on sequential days, a valuation of invasiveness could be obtained. Based on this method, we compared the invasiveness of the different cell types. Interestingly, although U87 cells showed the best rate of survival in our experiments (Fig. 4A), U251 cells were the most invasive cells. U251 cells showed the highest rate of surrounding tissue infiltration (206.7% ± 11.2%, n = 15) compared to HT1080 cells (138.8% ± 16.5%, p = 0.0107, n = 10) and U87 cells (165.9% ± 15.7%, p = 0.203, n = 8) (Fig. 4D). Importantly, the results observed in this study coincide with previous histopathological studies conducted in rodent models, which documented the differences in invasiveness between the two glioblastoma cell lines.51–53

Of note in this study, and of importance to measure invasiveness accurately, these measurements have to be done in 3D, as 2D projections of confocal stacks do not reflect the total invasiveness in all dimensions. Furthermore, cells need to be stably expressing the fluorophore as trials using dye labelled cells (CM-Di Dye) showed false positive rates of invasiveness due to detection of small fluorescent particles that may be either pure dye aggregates or vesicles released by tumor cells (Supplementary Fig. S2A, B). These small dye aggregates spread within the tissue in all dimensions, thus, including these particles leads to misleading results and an overestimation of invasiveness. Furthermore, an important prerequisite to measure invasiveness precisely is an additional labelling for microglia. Depending on the cell type, transplanted cells might be engulfed by microglia and signals will be detected within microglial phagosomes. These signals do not represent viable cells, and thus, do not reflect true infiltration. Only the additional microglial labelling allows the identification of these engulfed cells to exclude them from the analysis (Supplementary Fig. S2C, D).

Microglia promote U87 and U251 cell growth

To further investigate the direct role of microglia–glioma interactions in promoting survival, proliferation, and invasiveness of glioma cells, xenografting experiments were conducted in irf8−/− zebrafish that lack macrophages/microglia. Interferon regulatory factor 8 (irf8) is vital for macrophage development in mammals and in teleosts.47 The irf8 null mutant (irf8−/−) zebrafish was characterized to lack macrophages up to around 7 dpf and microglia were absent in the brain until 31 dpf.47 Thus we investigated xenografted U87, U251, and HT1080 cell survival and invasiveness in irf8−/− mutant zebrafish and compared the results to those observed in our mpeg1:EGFP xenograft studies.

Xenografts in irf8−/− zebrafish highlighted a strong impact of microglia on tumor progression for U87 cells and U251 cells. The results showed a significant reduction of U87 cell survival at all time points in irf8−/− mutants compared to mpeg1:EGFP (Fig. 5A). At 4 dpf the survival of U87 cells was 132% ± 16% (p < 0.0001) lower in irf8−/− mutants compared to mpeg1:EGFP (Fig. 5A). In line with this, we detected a significant decrease in tissue infiltration for U87 cells in irf8−/− zebrafish (96.9% ± 10.2%) versus mpeg1:EGFP (165.9% ± 15.7%) (Fig. 5D). The survival of U251 cells in irf8−/− zebrafish was also significantly reduced (Fig. 5B). At 4 dpf the survival of U251 cells was 41.4% ± 11.62% (p = 0.002) lower in irf8−/− mutants compared to mpeg1:EGFP (Fig. 5B). However, U251 tissue infiltration was not affected in irf8−/− zebrafish (195.5% ± 19.8%) compared to mpeg1:EGFP (206.7% ± 15.0%) (Fig. 5D).

Transplantation of HT1080 cells into irf8−/− zebrafish showed contrasting results. While HT1080 survival was rather poor in mpeg1:EGFP fish, a trend was detected toward better survival in the microglia mutant background (Fig. 5C). The results showed a nonsignificant increase of HT1080 cell survival by 27.1% ± 12.1% (p = 0.333) at 4 dpf in irf8−/− mutants compared to mpeg1:EGFP (Fig. 5C). Although better survival of HT1080 cells was observed in irf8−/− zebrafish, we did not detect an increase in tissue infiltration for HT1080 in the microglia mutant background (Fig. 5D).

To test if the protumoral function of the microglia can also be altered by pharmacological means in our xenograft model, we used the CSF-1R inhibitor BLZ945. CSF-1R signaling is crucial for survival of macrophages and microglia. BLZ945 has recently been shown to target microglia in a mouse pro-neural GBM model, leading to a regression of established tumors.26 To test BLZ945 in our model, we transplanted mpeg1:EGFP larvae with U87 cells at 3 dpf and incubated these larvae from 4 dpf (1 dpt) in 25 μM BLZ945 for 5 days. These larvae developed normally and did not show any adverse effects, confirming the high specificity for BLZ945 for the CSF-1R. Indeed, treatment with BLZ945 led to reduction of total microglial numbers in the optic tectum and to a decrease in U87 cell growth compared to controls (Fig. 6). Interestingly, as BLZ945 treatment did not lead to an ablation of all microglia, the impact on U87 cell growth was not as strong as observed before in the irf8−/− zebrafish (compare Figs. 5A and 6).

Together, these results clearly show that zebrafish microglia support the growth of human glioblastoma cells and, thus, further highlight the suitability of the zebrafish model to study human glioblastomas.

Discussion

The zebrafish as a model to study microglial responses toward glioma cells

The xenografting live-imaging assay used in this study presents a novel system that allows the response of the entire microglial network during glioma colonization to be studied in the living brain. Previous research has already demonstrated the power of the zebrafish model to study glioma growth. Several studies have transplanted glioma cells into the zebrafish yolk and shown that these cells form tumors, induce angiogenesis, and are responsive to treatment.34–36 Furthermore, intracranial models of glioma in larval zebrafish have been developed to study glioma invasiveness and angiogenesis37 and to use the zebrafish as a powerful model for drug screening.38,39 However, none of these studies have investigated the microglial response to xenografted cells. Thus we have established a larval zebrafish model to study microglia–glioma interactions in real time in the living brain.

We show that transplantation of U87, U251, and HT1080 cells into the optic tectum of larval zebrafish stimulated a microglia response that resulted in the recruitment of microglia cell bodies to the site of transplantation. The injection itself is minimally invasive, and media only with sham injections show a short injury response by microglial cells in direct proximity to the injection site. This microglial response is usually resolved within a few hours. However, the interactions observed upon
FIG. 5. Zebrafish microglia promote survival of U87 and U251 cells. U87, U251, and HT1080 cells were transplanted into the larval zebrafish tectum of mpeg1:EGFP and irf8\(^{-/-}\) zebrafish at 3 dpf and larvae were imaged at 0–4 dpt. (A) Comparison of survival for U87 cells in mpeg1:EGFP and irf8\(^{-/-}\) zebrafish over 4 days displayed as the ratio (expressed as a percentage) between the cell number at each time point and the starting cell number. (B) Comparison of survival for U251 cells in mpeg1:EGFP and irf8\(^{-/-}\) zebrafish over 4 days displayed as the ratio (expressed as a percentage) between the cell number at each time point and the starting cell number. (C) Comparison of survival for HT1080 cells in mpeg1:EGFP and irf8\(^{-/-}\) zebrafish over 4 days displayed as the ratio (expressed as a percentage) between the cell number at each time point and the starting cell number. (D) Comparison of the mean of cell invasiveness in mpeg1:EGFP and irf8\(^{-/-}\) zebrafish. \(n\) numbers represent the number of fish. All error bars represent the standard error of the mean. \(p\) values are indicated on graph (ns = nonsignificant).

FIG. 6. Inhibition of CSF-1R signaling leads to reduced microglia numbers and impairs U87 cell growth. U87 cells were transplanted into the larval zebrafish tectum of mpeg1:EGFP zebrafish at 3 dpf. Transplanted larvae were treated with 1% DMSO or 1% DMSO/25 \(\mu\)M BLZ945 from 4 dpf for 5 days. Microglial cell counts and U87 cell counts were done at 4 dpf before the start of treatment and the end of treatment (9 dpf). (A) Comparison of microglial cell numbers in control (1% DMSO) and BLZ945-treated zebrafish. Microglia numbers are displayed as the ratio (expressed as a percentage) of the final numbers at 9 dpf divided by the starting number at 4 dpf. (B) Comparison of U87 cell numbers in control (1% DMSO) and BLZ945-treated zebrafish. U87 cell numbers are displayed as the ratio (expressed as a percentage) of the final numbers at 9 dpf divided by the starting number at 4 dpf. All error bars represent the standard error of the mean. \(p\) values are indicated on graph.
flammation. These signals and others might lead to a pro-colony-stimulating factor expression and, thus, to trigger expression levels of IL-1β. The activation will vary depending on the molecular glioblastoma subtype. Interestingly, differential expression levels of immune genes have been detected between these glioblastoma subtypes. Interestingly, for U87 and U251 glioma cells, we observed microglia infiltration of the U251 cell mass and intimately interacted with U251 cells over the full duration of the experiment. Furthermore, in U251 samples, microglia were often seen to interact with U251 protrusions in the deeper tissue outside of the glioma central mass. In comparison, U87 cells attracted fewer microglia and these microglia showed only superficial interactions with U87 cells that diminished over time. These interactions were presented by individual microglial cells, which extended and retracted their protrusions to make surface contacts with the U87 cells. The differences in microglial responses toward U87 cells and U251 cells highlight intrinsic differences between these cell types. U251 cells are mutant for the tumor suppressor p53 and show high levels of expression for GFAP, vimentin, and S100β, for example. In contrast, U87 cells express wild-type tumor suppressor p53 and lack GFAP or S100β expression. Furthermore, U251 cells have been shown to express high levels of CCL2, which stimulates microglial responses. The low levels of CCL2 in U87 cells might be one of the reasons for the lower number of interactions with microglia cells observed in this study.

Our new model will facilitate new studies to reveal if microglia respond differently depending on glioblastoma subtype. Based on their gene expression, glioblastomas are classified into the Proneural, Neural, Mesenchymal, or Classical subtype. Interestingly, differential expression levels of immune genes have been detected between these glioblastoma subtypes. Thus, it is tempting to speculate that microglia recruitment and activation will vary depending on the molecular glioblastoma subtype.

Zebrfish microglia promote survival and invasiveness of human glioblastoma cells

U87 and U251 cells showed differences in growth and invasiveness upon transplantation into the zebrafish tectum. These differences are in line with differences in their in vitro growth rates and differences in invasiveness upon transplantation in rodent models. When investigating how zebrafish microglia activity affected these attributes, U87 and U251 glioblastoma cells were clearly influenced by microglia. This was demonstrated by the significant decrease in survival and invasiveness of U87 cells in the irf8a−/− zebrafish. In addition, U251 cell survival was also significantly decreased in the irf8a−/− zebrafish. Furthermore, treatment with the CSF-1R inhibitor BLZ945 resulted in decreased U87 cell growth as previously observed in a mouse proneural GBM model.

In summary, these experiments show that zebrafish microglia develop a protumoral activity and promote the growth of human glioblastoma cells. This highlights the suitability of the larval zebrafish model to investigate the role of microglia during brain tumor growth.

Conclusion

Finally, the fact that microglia respond differently depending on the glioma cell type could be very important when developing novel immunotherapeutics to target gliomas. Immunotherapeutics that specifically target microglia interactions may work on certain subtypes of gliomas, but the same desirable effect may not be achieved on others. The zebrafish xenograft assay presented in this study will become a powerful tool in cancer research to understand microglia–glioma interactions in detail in vivo. Due to advances in vertebrate animal screening technologies, zebrafish larvae can be screened and imaged automatically in 96-well plate format. Thus, based on our xenograft live imaging model, drug screening can be performed on a medium to high throughput level to identify drugs that impair the protumoral functions of microglia within gliomas.

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Disclosure Statement

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