Myeloid-Derived Suppressive Cells Promote B cell–Mediated Immunosuppression via Transfer of PD-L1 in Glioblastoma

Catalina Lee-Chang1, Aida Rashidi1, Jason Miska1, Peng Zhang1, Katarzyna C. Pituch1, David Hou1, Ting Xiao1, Mariafausta Fischietti2,3, Seong Jae Kang1, Christina L. Appin4, Craig Horbinski1,4, Leonidas C. Platanias2,3,5, Aurora Lopez-Rosas1, Yu Han1, Irina V. Balyasnikova1, and Maciej S. Lesniak1

Abstract

The potent immunosuppression induced by glioblastoma (GBM) is one of the primary obstacles to finding effective immunotherapies. One hallmark of the GBM-associated immunosuppressive landscape is the massive infiltration of myeloid-derived suppressor cells (MDSC) and, to a lesser extent, regulatory T cells (Treg) within the tumor microenvironment. Here, we showed that regulatory B cells (Breg) are a prominent feature of the GBM microenvironment in both preclinical models and clinical samples. Forty percent of GBM patients (n = 60) scored positive for B-cell tumor infiltration. Human and mouse GBM-associated Bregs were characterized by immunosuppressive toward activated CD8+ T cells, the overexpression of inhibitory molecules PD-L1 and CD155, and production of immunosuppressive cytokines TGFβ and IL10. Local delivery of B cell–depleting anti-CD20 immunotherapy improved overall survival of animals (IgG vs. anti-CD20 mean survival: 18.5 vs. 33 days, P = 0.0001), suggesting a potential role of Bregs in GBM progression. We unveiled that GBM-associated MDSCs promoted regulatory B-cell function by delivering microvesicles transporting membrane-bound PD-L1, able to be up-taken by tumor B cells. The transfer of functional PD-L1 via microvesicles conferred Bregs the potential to suppress CD8+ T-cell activation and acquisition of an effector phenotype. This work uncovered the role of B cells in GBM physiopathology and provides a mechanism by which the GBM microenvironment controls B cell–mediated immunosuppression.

See related Spotlight on p. 1902

Introduction

Glioblastoma (GBM), an incurable malignant brain cancer, is characterized by its ability to build an immunosuppressive environment, which blocks antitumor immunity and promotes tumorigenesis. There are many drivers of GBM-induced immunosuppression, including the production of immunoregulatory secreted factors such as cytokines (TGFβ and IL10), enzymes (IDO), or prostaglandins (1–3). Cellular components such as regulatory T cells (Treg), myeloid-derived suppressor cells (MDSC), and tumor-associated macrophages (TAM) are also known to mediate GBM-associated immunosuppression (4–6). MDSCs are known to act as major immunosuppressive cells within the tumor vicinity. They have also been found in the peripheral blood of GBM patients, suggesting a systemic influence on immunity (5). Inhibitory molecules expressed on the surface of GBM and myeloid suppressive cells, such as programmed death ligand 1 (PD-L1) or CD155 (poliovirus receptor), are known to inhibit the effector function of T cells (7, 8), and it has been suggested that the GBM-derived hypoxic environment might contribute to MDSC and Treg function (9, 10).

Although immunotherapy remains a promising treatment for GBM (11), a better understanding of how GBM controls immunity to promote its survival, progression, and immune escape is necessary to optimize current immunotherapeutic approaches. Our group and others have reported the importance of GBM-associated immunosuppressive factors such as MDSCs or, to a lesser extent, Tregs as fundamental immunosuppressive players in GBM (12, 13). Other players of the immune system, such as B cells, have also been reported to be part of the GBM immune landscape (14). However, whether tumor-infiltrating B cells play a relevant role in GBM progression remains to be elucidated.

This present work provides a comprehensive phenotype and functional profile of GBM regulatory B cells (Breg). We showed that human and murine GBM-associated B cells exhibit an immunosuppressive phenotype characterized by the presence of inhibitory molecules PD-L1 and CD155 and production of immunoregulatory cytokines TGFβ and IL10. Bregs represented ~10% of bone marrow–derived infiltrating immune cells in two different
orthotopic brain tumor models (GL261 and CT2A), and 40% of GBM patients who were screened scored positive for B-cell tumor infiltration. GBM-associated B cells showed an immunosuppressive function toward activated CD8^+ T cells, and their pathophysiologic relevance was highlighted by extended animal survival after local delivery of B cell–depleting immunotherapy. We showed that the tumor microenvironment, and more precisely, MDSCs, played a fundamental role in promoting immunosuppressive B cells. MDSCs mediated transfer of membrane-bound PD-L1 to B cells, resulting in the promotion of B cell–mediated immunosuppression. Overall, the present study provides an extensive characterization of GBM-associated B cells and unveiled a mechanism of intercellular communication between MDSCs and B cells to expand MDSC immunosuppressive effects.

**Materials and Methods**

**Human samples**

All human samples (tumor and peripheral blood) and frozen tissue were collected by the Nervous System Tumor Bank of the Northwestern University. Human samples were collected in EDTA tubes. Peripheral blood samples from GBM patients were collected in EDTA tubes. Peripheral blood mononuclear cells (PBMCs) were isolated using the Ficoll (GE Healthcare) gradient. Tumor cells and PBMCs were immediately set in complete RPMI medium [RPMI + 10% heat-inactivated fetal bovine serum (FBS), 10 mmol/L HEPES–sodium pyruvate, 1 mmol/L sodium pyruvate, 0.01% 2-mercaptoethanol, 2 mmol/L L-glutamine, penicillin (100 U/mL), and streptomycin (100 μg/mL); all reagents from Thermo Fisher].

**GBM B cell–mediated T-cell suppression assay**

The assay was performed in an autologous manner, and thus, B cells (from tumor and PBMCs) and T cells (PBMCs) were from the same patient. B cells from tumor and PBMCs were obtained using the EasySep Human CD19 Positive Selection Kit II (STEMCELL Technologies). PBMC T cells were isolated using the EasySep Human T-Cell Isolation Kit (STEMCELL Technologies) and labeled with 10 μmol/L of the eBioscience cell proliferation dye eFluor 450 (Thermo Fisher). Cells were activated with T-cell activator anti-CD3/CD28 beads (Dynabeads, Invitrogen, Thermo Fisher) at 1:3 beads:T-cell ratio supplemented with IL2 (50 μ/mL, PeproTech) and cocultured at a 1:1 ratio with tumor-infiltrating or PBMC CD19^+ B cells for 72 hours. CD4^+ and CD8^+ T-cell proliferation (eFluor 450 dilution) and activation status [intracellular granzyme B (GzmB) and IFNγ expression] were analyzed using flow cytometry (Supplementary Table S1).

**Tumor-infiltrating CD163^+ cell isolation and microvesicle uptake**

Tumor cells and PBMCs were obtained as described above, and CD163^-+ macrophages were isolated using an anti-human CD163 biotin (clone GHI/61, BioLegend) and the anti-biotin Microbeads (Miltenyi Biotec). Cells were magnetically isolated using LS columns (Miltenyi Biotec). CD163^-+ cells were labeled with the lipophilic dye CellTrace Violet (CITV, Invitrogen, Thermo Fisher) and placed in the upper chamber of a 0.4-μm transwell system in complete RPMI. In the lower chamber, PBMCs from the same donor were placed at 10^6 cells/mL in complete RPMI. After 24 hours, cells from the bottom chamber were harvested and tested by flow cytometry the acquisition of the CTV dye by B cells, CD4^- Foxp3^- Tregs, and CD33^- myeloid cells by flow cytometry. See Supplementary Table S2 for antibody information.

**Mice**

C57BL/6, CD45.1 C57BL/6, B cell–deficient (μMT, BKO), and IL10-deficient (IL10 KO) mice were from The Jackson Laboratory. Animals were 6 to 8 weeks old at the time of the experiment initiation. All animal experimentation protocols are approved by the Institutional Animal Care and Use Committee (IACUC) under the protocol # IS00002459 at the Northwestern University. All animals were housed in specific pathogen–free animal facility at Northwestern University.

**Cell lines**

GL261 cells were obtained from the National Cancer Institute (NCI), and CT2A cells were a gift from Pr. Tom Seyfried (Boston College). The GL261 cell line identity and purity were evaluated.
Brain tumor injection

A total of $2 \times 10^3$ GL261 or CT2A cells were intracranially (i.c.) implanted as previously described (2). Mice were anesthetized through intraperitoneal administration of a stock solution containing ketamine (100 mg/kg) and xylazine (10 mg/kg). The surgical site was shaved and prepared with a swab of povidone-iodine followed by a 70% ethanol. The shaving procedure was performed three times in total. An incision was made at the midline for access to the skull. A 1-mm-diameter burr hole was drilled 2 mm posterior to the coronal suture and 2 mm lateral to the sagittal suture. Mice were then placed in a stereotactic frame, and tumor cells were injected in a total volume of 2.5 μL using a Hamilton syringe fitted with a 26-gauge blunt needle at a depth of 3 mm. The incision was then stapled closed.

Cannula implantation

Briefly, mice were anesthetized, and a skin incision ~10 mm in length was made over the middle frontal to the parietal bone to expose the surface of the skull. A 26-gauge sterile guide cannula for mice (Plastics One) was installed into the mouse brain at 2 mm depth through the burr hole generated during tumor implantation as described above. Tissue glue was applied around the burr hole to secure the protrusion of the cannula for long-term stable positioning. The scalp was closed with surgical glue around the hole to secure the protrusion of the cannula for long-term stable positioning. If endpoint was reached, mice were euthanized. For intracranial administration of CD20-depleting Ab was performed using a cannula implantation system 7 and 10 days after tumor implantation. Briefly, a 33-gauge sterile syringe was inserted into the guide cannula. The syringe was covered with a sleeve designed to extend 1 mm beyond the tip of the guide cannula. The diluted anti-CD20 in sterile 0.9% saline was injected into the brain (25 μg/mouse/injection; final volume of 2.5μl/injection). After injection, the cannula was covered using a 33-gauge dummy cannula for mice.

Brain, lymphoid tissue, and blood single-cell suspensions

Mice were bled retro-orbitally, and blood samples were collected in heparinized-PBS solution (1 mg/mL, Sigma-Aldrich). Red blood cells were lysed using an ACK lysis solution (Gibco, Thermo Fisher). After blood collection, mice were euthanized in a CO₂ chamber and intracardially perfused with chilled PBS. Brain single-cell suspensions were obtained by mechanical dissociation using a manual tissue homogenizer (Potter-Elvehjem PTFE pestle, Sigma-Aldrich) in HBSS. Myelin and debris were removed by Percoll gradient separation. Leukocytes from deep and superficial cerebral lymph nodes were obtained by mechanical tissue dissociated using a 70-μm cell strainer and a syringe plunger. Brain, blood, and lymph node cells were collected in complete RPMI media. Cells were used for immunophenotype analysis or ex vivo functional assays as described below.

Flow cytometry and immunophenotype analysis

Immunophenotype analysis of immune cells from tumor-bearing mice was performed at different time points (0, 7, 14, and 21 days after tumor injection) as described below. In some experiments, the immunophenotype analysis was performed at one time point defined in the Results section. After collection single-cell suspensions, cells were counted and washed with staining buffer (5% bovine serum albumin, 0.001% sodium azide in PBS). Cells were incubated with 1 μL Fc receptor blocking Ab (anti-CD16/32, clone 93, BioLegend) per 10^6 cells in 100 μL staining buffer for 5 minutes at room temperature. For surface staining, cells were incubated with 1 μL Ab per 10^6 cells for 30 minutes at 4°C. Cells were washed twice with cold PBS. Cells were stained with Fixable Viability Dye eFluor 780 (eBioscience, Thermo Fisher) for 30 minutes at 4°C. Cells were washed twice with staining buffer. For intracellular staining, cells were fixed and permeabilized using the eBioscience Foxp3/Transcription Factor Staining Buffer Set (Innogenot, Thermo Fisher) for 90 minutes at room temperature. Cells were washed twice with the Permeabilization Buffer (provided in the permeabilization/fixation buffer kit) and incubated with 1 μL Ab for 1 hour at 4°C. Cells were washed twice with staining buffer. To evaluate cytokine expression, cells were stimulated for 5 hours at 37°C with the eBioscience Cell Stimulation Cocktail plus protein transport inhibitors (500×, Thermo Fisher) prior staining. The list of Abs used to analyze the different immune cells population can be found in Supplementary Tables S1, S3, and S4. All Abs were from BioLegend. Data were acquired with BD FACSDroped Symphony analyzer and analyzed with FlowJo software. Dead cells and debris were excluded using the Live/Dead staining (Viability Dye eFluor 780). B cells were identified as CD45^− CD11b^− CD19^+ cells. CD4^+ and CD8^+ T cells were identified as CD45^+ CD11b^− CD4^+ or CD45^+ CD11b^− CD8^+ cells, respectively. Monocytic MDSCs were
identified as CD45+CD11b+CD11c-Ly6C+Ly6Glow cells. Polymorphonuclear (PMN) MDSCs were identified as CD45+CD11b+CD11c-Ly6C-Ly6G+ cells. All Ab details are found in Supplementary Tables S1, S3, and S4.

Human B-cell phenotype was assessed from tumor cells and PBMCs obtained as described above. Cells were processed using the same method as for murine cells. B cells were defined as CD45+CD11b-CD19+CD20+ cells, and expression of PD-L1, CD155, II.10, and LAP (TGFβ) were assessed. All Abs were from BioLegend unless otherwise specified. Details are found in Supplementary Table S2.

**In vitro murine MDSC generation**

GBM-associated MDSCs were generated as in previous reports (13, 15). Bone marrow (BM) cells from C57BL/6 mice were flushed out from femurs with complete RPMI using a 10 ml syringe and 25-gauge needle into complete RPMI (Corning). BM cells were centrifuged (10 minutes, 1,500 RPM, at 4°C), and red blood cells were lysed using ACK lysis buffer (Sigma) for 5 minutes at RT. Cells were washed (5 minutes, 1,500 RPM, at 4°C) with complete RPMI, counted, and plated into 24-well plates (Corning) at a density of 2×10^6 cells/well with 50% complete RPMI, 50% conditioned media (CM; 8 ml DMEM (Corning) supernatant that was collected from original seeding of 2×10^6 CT2A cells after 72 hours of culture), and GM-CSF (40 ng/ml; PeproTech). After 3 days of culture, old media were removed by aspiration, and new media (same as aforementioned) were added to the cells for an additional 3 days of incubation. Six days after BM isolation, cells were lifted by pipetting up and down, washed (10 minutes, 500 × g, at 4°C), and stained for characteristic flow-cytometric analysis (CD11b+Ly6C+PD-L1+arginase-1+; Supplementary Table S3) and further functional assays as described below.

**In vitro human MDSC generation**

Human monocytes were isolated from buffy coat samples using the Human Monocyte Isolation Kit (STEMCELL Technologies). Cells were counted and plated in a 24-well plate with the density of 10^6 per well with 50% complete RPMI (20% FBS), 50% CM (8 ml DMEM (2% FBS) supernatant that was collected from original seeding of 2×10^6 GBM6, GBM43, GM12, and MES83 cells, after confluence of the cells, human GM-CSF (80 ng/ml), and human IL6 (80 ng/ml; PeproTech). After 3 days of culture, old media were removed and replaced with new media (same as aforementioned) on days 3 and 6 of culture. After 9 days of culture, cells were lifted, washed (5 minutes, 1,500 RPM at RT), and tested for the expression of arginase-1 and PD-L1 by flow cytometry (Supplementary Table S2).

**Murine and human Breg conversion assay by MDSCs**

Splenic B cells from naive mice were isolated using the EasySep Mouse B-cell Isolation Kit (STEMCELL Technologies). To test the ability of MDSCs to convert naïve B cells into Bregs, cells were cultured for 48 hours at a 1:1 ratio. Then, B cells were analyzed for their expression of PD-L1, CD155, TGFβ (LAP), and II.10 (Supplementary Table S3 for Ab details). Alternatively, cells were cultured in a 0.4-μm membrane transwell system (Corning, Life Sciences). MDSCs were placed in the upper compartment and naïve B cells supplemented with 100 mmol/L murine BAFF (BioLegend) in the lower compartment, at a 1:1 ratio. Coculture was maintained for 12, 24, 48, or 72 hours unless otherwise specified. For human Breg conversion using human MDSCs, B cells were isolated using the EasySep Human B-cell Isolation Kit (STEMCELL Technologies). B cells were incubated with 100 mmol/L recombinant human BAFF (PeproTech).

**In vivo BLZ954 treatment and Breg conversion**

B cell–deficient mice (μMT, BKO) were injected intracranially with 2×10^5 CT2A cells, as described above. The CSF1-R inhibitor BLZ954 (Novartis) was dissolved in Captisol and administered daily by gavage (200 mg/kg) from days 3 to 7 after tumor inoculation. B cells from CD45.1+ C57BL/6 mouse spleens were isolated using the EasySep Mouse B-cell Isolation Kit (STEMCELL Technologies, purity >97%) in injected intravenously (retro-orbital injection) 24 hours after BLZ954 treatment termination. Forty-eight hours after, CD45.1+ B cells (clone A20, BioLegend) were assessed by flow cytometry for their expression of CD155 and PD-L1 (Supplementary Table S4 for Ab details).

**In vivo PD-L1 blockade and Breg conversion**

B cell–deficient mice (μMT, BKO) were injected intracranially with 2×10^5 CT2A cells. Mouse brains were irradiated with 3 Gy for 3 consecutive days (days 8–10) using a Gammacell 40 Exactor (Best Theratronics). Three days later, mice received 5 μL of blocking anti–PD-L1 (clone 10F.9G2, Bio X Cell, 5 μg/injection) intracranially via cannula. Two days later, mice received intracranially 5×10^5 splenic B cells, negatively isolated from CD45.1+ C57BL/6 spleens using the EasySep Mouse B-cell Isolation Kit (STEMCELL Technologies, purity >97%). Forty-eight hours after the adoptive transfer, mice were sacrificed, and B cells were isolated from tumor-bearing brains using the EasySep Mouse CD19 Positive Selection Kit II (STEMCELL Technologies). B-cell purity (≥96%) was evaluated by the congeneric marker CD45.1 (clone A20, BioLegend). B cells were tested for their suppressive function against activated splenic CD8+ T cells as described below.

**Murine CD8+ T-cell suppression assay**

CD8+ T cells were isolated from spleens of naïve mice using EasySep Mouse CD8+ T-Cell Isolation Kit (STEMCELL Technologies). Cells were labeled with the eBioscience cell proliferation dye eFluor 450 (Thermo Fisher). To test B-cell ability to suppress activated CD8+ T cells, proliferation, cells were mixed at a 1:1 BT ratio. CD8+ T-cell activation was assessed using the anti-CD3/CD28 T-cell activating beads used at 1:3 beads:CD8+ T cells (Invitrogen, Thermo Fisher) for 4 days in complete RPMI. T-cell proliferation (eFluor 450 dye dilution) and effector T-cell factors such as GzmB and IFNγ were assessed by flow cytometry. Anti-II.10 (Thermo Fisher), anti-TGFβ (R&D Systems), anti-TIGIT (BioLegend), and anti-PD-L1 (Bio X Cell) were added every day in the culture at 10 μg/mL. Alternatively, expression of TGFβ (LAP), II.10, and IFNγ by B cells while cocultured with CD8+ T cells was examined after 36 hours (Supplementary Table S4).

**MDSC-derived microvesicle isolation, quantification, and labeling uptake by B cells**

Microvesicles (MW) were prepared from MDSC CM as previously described (16, 17). Briefly, MDSC-CM was collected after initial centrifugation to pellet life cells at 300 × g, 5 minutes at room temperature. Supernatants were cleared of dead cells, cell
debris, and large vesicles by centrifugation at 2000 \times g for 30 minutes at 4°C. Apoptotic bodies were removed by ultracentrifugation at 10,000 \times g for 30 minutes at 4°C. Remaining supernatant was ultracentrifuged at 100,000 \times g for 90 minutes at 4°C. Pellets containing vesicles were resuspended in cold PBS and ultracentrifuged at 100,000 \times g for 90 minutes at 4°C. MV pellets were resuspended in 50 to 100 \mu L of ice-cold PBS and used for a downstream application. The particle size distribution of MVs was measured by dynamic light scattering (DLS) using a Zetasizer Nano ZSP (Malvern Panalytical) and presented as diameter in nm and polydispersity index. The surface charge of MV was determined by zeta-potential using a Zetasizer Nano ZSP (Malvern Panalytical).

To label MDSC-derived MVs, MDSCs were labeled with the lipophilic dye CTV (Invitrogen) for 5 minutes at room temperature. Dye excess was removed by washing cells 3 times with complete RPMI. MVs were isolated from supernatants as described above. To test the MV uptake by B cells, CTV-labeled MDSC were cocultured with splenic B cells isolated from naïve mice using the EasySep Mouse B-cell Isolation Kit (STEMCELL Technologies) in a 0.4-μm transwell system (Corning, Life Sciences). B cells were harvested at different time points depending on the experiment and evaluated for the acquisition of the CTV fluorescence by flow cytometry. In some experiments, MDSCs were incubated for 2 hours at 37°C with the MV release inhibitor GW4869 (20 μmol/L, Cayman Chemicals). Cells were washed 3 times with complete RPMI and plated at 1 × 10^6 cells/mL with B cells. Alternatively, MVs were fluorescently labeled with PKH67 green fluorescent cell linker kit (Sigma-Aldrich) following the manufacturer's instructions. Briefly, 1 μL of PKH67 dyes in 150 μL of Diluent C solution were mixed with the equal volume of MV (from 10^6 cells/mL) at room temperature for 5 minutes. To remove unlabelled PKH67 dyes, 1.5 mL of 0.971 M sucrose solution was carefully added into the bottom of the tube, following which 8 mL of serum-free media was added to PKH67-labeled MVs. After centrifugation at 200,000 \times g for 2 hours, unlabelled PKH67 dyes located in the interface layer were aspirated. PKH67-labeled MVs located in the bottom were resuspended with PBS. MV content was normalized by their protein content using a standard Bradford method before incubation with B cells. A range of 5 to 15 μg of protein was used across the experiments to convert B cells into Bregs. However, the same protein content was used among groups within the same experiment.

**PD-L1 knockdown using siRNA**

Lipid nanoparticles (LNP) for siRNA transfection were formulated by 1,2-dioleoyl-3-trimethylammonium-propane, cholesterol, and 1,2-distearoyl-sn-glycero-3-phosphoethanolamine-N-[methoxy(polyethylene glycol)-2000] (Avanti Polar Lipids), and synthesized using a thin-film rehydration and sonication method. pd-L1 siRNA (Sigma) was encapsulated into LNP by coinoculation in HEPES buffer for 30 minutes at room temperature. MDSCs were lifted and replated at 10^6 cells/mL/well in 24-well plates and allowed to adhere overnight. MDSCs were incubated with siRNA (pd-L1 or scramble)/LNP complex in Opti-MEM at siRNA concentration of 100 nmol/L for 4 hours at 37°C. The medium was replaced by complete RPMI, and the cells were cultured for 24 hours at 37°C. RNA was isolated from siRNA-treated MDSCs using the RNEasy Plus Mini Kit (QIagen). Following chloroform extraction, samples were loaded onto RNEasy Plus columns (QIagen) and processed according to the manufacturer’s protocol. Total RNA was quantified using a Nanodrop apparatus (Thermo Scientific) and then converted to cDNA using the iScript cDNA synthesis kit (Bio-Rad). pd-L1 and control β-actin transcripts were analyzed by using the 2^ΔΔCT method. Murine pd-L1 primers: forward TGCTGCTATAACGCTAACC; reverse CCACGAAATTCTCTGGTTG (18). Murine β-actin primers: forward TTGTCGA-CAGAGTGCAGAAAG; reverse ACACTGCTGGAAAGTGAGAC (19).

**Membrane–cytosol fractionation and immunoblotting**

MDSCs and MVs were lysed using the Mem-PER Plus Kit (Thermo Scientific, #89842) according to the manufacturer's instructions. A total of 10 μg of protein/samples/fraction was resolved by SDS-PAGE (Bio-Rad), transferred to Immobilon-P PVDF membranes (Millipore), then probed with primary Abs anti-CD45 (1/1,000, clone 72787, Cell Signaling Technology), anti-PD-L1 (1/1,000, clone BE1010, Bio X Cell), and anti–GAPDH (1/15,000, clone MAB374, Millipore). The GE Healthcare Amersham ECL anti-rabbit-HRP (Thermo Fisher), and the Goat Anti-Mouse IgG (H + L)-HRP (Bio-Rad) were used as secondary Abs at 1:5,000 dilution. After reaction with Amersham ECL detection reagent (GE Healthcare), blots were visualized for the indicated proteins using autoradiography.

**Cryoelectronic microscopy**

MV were imaged by cryoelectron microscopy using the software package Digital Micrograph (Gatan, Inc.) using a JEOL 3200FS Transmission Electron Microscope (JEOL USA, Inc.) equipped with an in-column energy filter (omega filter) operating at 300 kV at a magnification of 30,000×. Images were recorded using a K2 Summit Direct Electron Detection camera (Gatan, Inc.) in counting mode, and motion corrected using tools in Digital Micrograph. Sample preparation utilized 400 mesh lacey Carbon copper grids (LC400-CU, Electron Microscopy Sciences), glow discharged at either 5 watts, 10 watts, or 25 watts for 10 seconds, with a Pelco easiGlow 91000 Glow Discharge Cleaning System (Ted Pella, Inc.). Grids were vitrified by plunging into liquid ethane. Sample (4 μL) was applied to each prepared grid and incubated for 30 seconds under ≥95% humidity at 4°C using a Vitrobot Mark IV cryo system (FEI/Thermo Fisher). Blot forces of 0.0 and 1.0 and a range of blot times, from 1 to 4 seconds were tested.

**MV uptake inhibition assay**

MDSCs were incubated for 2 hours with endocytosis inhibitors. Supplementary Table S5 describes the type of inhibitor and the concentration used. MVs were derived from MDSCs fluorescently labeled with the lipophilic dye (CTV) described above. Cells were then incubated with MVs (10^6 cells/mL) at room temperature for 5 minutes. After centrifugation with PBS, MV content was normalized by their protein content using a standard Bradford method before incubation with B cells. A range of 5 to 15 μg of protein was used across the experiments to convert B cells into Bregs. However, the same protein content was used among groups within the same experiment.

**Statistical analysis**

Data are shown as mean ± SD for a continuous variable and number (percentage) for a categorical variable. Differences between two groups were analyzed by Student t test or Wilcoxon signed-rank test.
rank-sum test as appropriate. Differences among multiple groups were evaluated using one-way ANOVA with post hoc Tukey test, or Kruskal–Wallis H Tests followed by post hoc Dunn multiple tests as appropriate. Survival curves were generated via the Kaplan–Meier method and compared by log-rank test and multiple comparisons were adjusted using Bonferroni method. Categorical variables were analyzed using Fisher exact tests or $\chi^2$ tests as appropriate. All the tests are two-sided, and $P$ values or Benjamini–Hochberg adjusted false discovery rates less than 0.05 were considered as significant. Statistical analyses were performed using SAS9.4 and GraphPad Prism7.03.

**Results**

**Functional and immunophenotypic characterization of GBM-infiltrating B cells**

We first evaluated whether infiltration of B cells occurred in tumors of GBM patients. A total of 60 GBM tissue sections were analyzed for the expression of CD20. Forty percent of the samples scored positive for CD20, and B-cell infiltration showed a predominant perivascular distribution (Table 1). Our data showed that B-cell recruitment did not significantly correlate with age or gender of the patient, nor the IDH mutation or MGMT methylation status of the tumor (Table 1). CD20 gene $(M S 4 A 1)\text{ expression was found higher in grade IV gliomas only when compared with grade II gliomas (P} = 0.05;\text{ Supplementary Fig. S1A; TCGA database). These data supported the lack of correlation between GBM patient survival and CD20 gene expression in tumors (Supplementary Fig. S1B; TCGA database). In the context of GBM, B cells have been reported to exhibit immunosuppressive function toward activated CD8$^+$ T cells (20, 21) in vitro, suggesting a potential role in protumorigenic immune responses. We observed that B cells isolated from freshly resected tumors suppressed activated autologous CD8$^+$ T-cell proliferation and expression of the cytotoxic factor GzmB (Fig. 1A). No significant effect was observed in CD4$^+$ T-cell activation (Supplementary Fig. S1C). Our histopathologic evaluation showed that B-cell infiltrates were in close proximity to CD8$^+$ T cells in the perivascular areas of the tumor (Fig. 1B; Table 1).

The analysis of the immune landscape in GBM orthotopic GL261 and CT2A experimental models revealed that B cells infiltrated the tumor. Absolute counts of B cells in the brain were significantly increased after tumor implantation (Supplementary Fig. S1D and S1E). The presence of B cells in the tumor correlated with an overall B-cell lymphopenia in the deep cervical lymph nodes (dCLN) and circulation (Supplementary Fig. S1D and S1E). B cells from the brain, blood, dCLN, and superficial CLN from either CT2A or GL261 tumor-bearing mice were isolated and tested for their ability to suppress splenic CD8$^+$ T-cell activation (Fig. 1C; Supplementary Fig. S2A). Activated CD8$^+$ T cells without B cells (No B Act) were used as a positive control for proliferation. B cells from the dCLN of naïve mice were used as an internal control as our data showed that these B cells functioned as an enhancer of CD8$^+$ T-cell activation (dCLN B; Fig. 1C). Tumor-infiltrating B cells from CT2A (Fig. 1C and D) or GL261 (Supplementary Fig. S2A) significantly suppressed activated CD8$^+$ T-cell proliferation. Circulating B cells and B cells from the dCLN and the superficial CLN from both CT2A- and GL261-bearing mice did not affect CD8$^+$ T-cell expansion, suggesting that B cells’ suppressive function is a unique feature of the B cells located in the tumor microenvironment. Tumor-associated B cells also inhibited the acquisition of effector phenotype by CD8$^+$ T cells, as seen by the inhibition of GzmB and IFN$\gamma$ expression (Fig. 1D; Supplementary Fig. S2A). These data showed that GBM-infiltrating B cells had the capability to suppress proliferation and expansion of effector CD8$^+$ T cells.

To evaluate the possible mechanisms controlling B cell–mediated CD8$^+$ T-cell suppression, we analyzed the immunophenotype of GBM-associated B cells. Tumor-infiltrating B cells from 5 different GBM patients showed overexpression of inhibitory molecules PD-L1, the poliovirus receptor (PVR) CD155, and immunosuppressive cytokines IL10 and TGF$\beta$ (LAP; Fig. 2A). The GBM-associated immunosuppressive phenotype observed in humans was also found in both CT2A (Fig. 2B) and GL261 (Supplementary Fig. S2B) murine models. The overexpression of CD155, IL10, and TGF$\beta$ was predominant during the earlier stages of the tumor progression (days 7 and 14 after tumor cell injection), whereas PD-L1 expression was at its highest at late stages (days 14 and 21 after tumor implantation). Overall, our data suggested that B cells infiltrating the tumor vicinity were Bregs that could inhibit CD8$^+$ T-cell activation and acquisition of effector cytotoxic properties. The collection of molecules expressed by GBM-associated B cells is well known to play an important role in cancer-mediated immunosuppression and immunosurveillance escape and is under investigation as a potential target for the development of immunotherapies (22–28). Supporting this current knowledge, neutralization of soluble IL10 and TGF$\beta$, blockade of TIGIT (ligand for CD155) on CD8$^+$ T cells, and PD-L1 on

### Table 1. GBM patient characteristics and tumor analysis for B-cell infiltration

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<td>Female</td>
<td>6 (24%)</td>
<td>12 (32.43%)</td>
<td>–</td>
</tr>
<tr>
<td>New or recurrence</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>New</td>
<td>18 (72%)</td>
<td>25 (67.57%)</td>
<td>0.7840</td>
</tr>
<tr>
<td>Recurrence</td>
<td>7 (28%)</td>
<td>12 (32.43%)</td>
<td>–</td>
</tr>
<tr>
<td>IDH mutation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>5 (20%)</td>
<td>6 (16.22%)</td>
<td>–</td>
</tr>
<tr>
<td>No</td>
<td>20 (80%)</td>
<td>30 (81.08%)</td>
<td>–</td>
</tr>
<tr>
<td>Unknown</td>
<td>0 (0%)</td>
<td>1 (2.7%)</td>
<td>–</td>
</tr>
<tr>
<td>MGMT methylation</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Positive</td>
<td>9 (36%)</td>
<td>20 (54.05%)</td>
<td>0.4683</td>
</tr>
<tr>
<td>Negative</td>
<td>15 (60%)</td>
<td>16 (45.24%)</td>
<td>–</td>
</tr>
<tr>
<td>N/A</td>
<td>1 (4%)</td>
<td>1 (2.7%)</td>
<td>–</td>
</tr>
<tr>
<td>Associated with CD8</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Yes</td>
<td>25 (100%)</td>
<td>0 (0%)</td>
<td>&lt;0.0001</td>
</tr>
<tr>
<td>No</td>
<td>0 (0%)</td>
<td>37 (100%)</td>
<td>–</td>
</tr>
<tr>
<td>Location</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Perivascular</td>
<td>23 (92%)</td>
<td>0 (0%)</td>
<td>–</td>
</tr>
<tr>
<td>Perivascular intraparenchymal</td>
<td>2 (8%)</td>
<td>0 (0%)</td>
<td>–</td>
</tr>
</tbody>
</table>

**NOTE:** A total of 60 GBM patient tumor samples were tested for the presence of CD20$^+$ B cells by IHC. Patients were stratified (variable) by age, gender, newly diagnosed or recurrent, isocitrate dehydrogenase (IDH) mutation, and O(6)-methylguanine-DNA methyltransferase (MGMT) methylation status, CD8$^+$ T-cell infiltration, and location. These variables were analyzed for statistical correlation with the presence or not of B cells (P value).

**Abbreviation:** N/A, not applicable.
tumor-infiltrating B cells using inhibitory Abs rescued CD8<sup>+</sup> T-cell proliferation and expression of GzmB and IFN<sub>γ</sub> (Fig. 2C), with anti–PD-L1 being the most effective regimen. Taken together, our results suggested that GBM Bregs negatively regulated activation of CD8<sup>+</sup> T cells by expression of membrane inhibitory molecules PD-L1 and CD155 and production of immunosuppressive soluble factors such as IL10 and TGFβ.

**Anti-CD20–depleting immunotherapy provides an extended animal survival**

To evaluate the pathophysiologic relevance of GBM Bregs in tumor progression, we tested the effects of B cell–depleting immunotherapy using anti-CD20. Because the immunosuppressive function of GBM Bregs appeared to be restricted to the tumor and not the periphery, we aimed to preferentially deplete B cells in
To do so, we delivered the CD20-depleting Ab intracranially via cannula after 7 days of CT2A tumor implantation. We observed that animals receiving the B cell–depleting immunotherapy survived significantly longer than the IgG control group (IgG vs. anti-CD20 mean survival: 18.5 vs. 33 days, \( P = 0.0001 \); Fig. 3A). In an independent experiment, we confirmed local injections of B cell–depleting Ab only partially depleted peripheral B cells (blood, spleen, and dCLN; Supplementary Fig. S3A). The extended animal survival after local B-cell–depleting immunotherapy correlated with increased tumor-infiltrating effector CD8\(^+\) T cells expressing GzmB and IFN\(\gamma\) (Fig. 3B). However, no survival benefit was observed in the animals receiving the immunotherapy systemically (intraperitoneal injection; Supplementary Fig. S3B and S3C).

Identification of Regulatory B Cells in GBM

Figure 2.
Glioma-associated B-cell phenotype.
A, Five paired tumor and PBMC samples from newly diagnosed GBM patients were analyzed for the presence of B cells and their phenotype by flow cytometry. PD-L1, CD155, TGF\(\beta\) (LAP), and IL10 were assessed (\( n = 4 \) patients). Difference between tumor and PBMC B-cell phenotype was assessed by unpaired \( t \) test. B, CD155, PD-L1, IL10, and TGF\(\beta\) (LAP) expression was analyzed in B cells from tumor (brain), blood, dCLNs, and superficial CLNs at 7, 14, and 21 days after CT2A tumor implantation. \( n = 4 \) mice/group in two independent experiments. For comparisons of the mean differences between groups based on two factors (time and tissue), ordinary two-way ANOVA with Tukey multiple comparison test was used. C, B cells from CT2A tumors (\( n = 5/\)experiment) were tested for suppression of CD8\(^+\) T-cell activation after treatment with blocking anti-IL10, anti-TGF\(\beta\), anti-TIGIT, or anti-PD-L1 Abs were added every day throughout the experiment (72 hours). CD8\(^+\) T-cell activation was assessed by cell proliferation and expression of intracellular GzmB and IFN\(\gamma\). Representative experiment of three independent experiments performed in triplicates. All histograms mean \pm SD of triplicates. In all experiments: ns, not statistically significant; \(*\), \( P < 0.05 \); \(*\)*, \( P < 0.01 \); \(*\)***, \( P < 0.001 \); \(*\)***, \( P < 0.0001 \).
Supporting our observations using B-cell depletion, B cell–deficient μMT (B-cell KO) mice survived significantly longer than wild-type (WT) controls (Fig. 4A), and this correlated with increased expression of GzmB by CD8+ T cells (Supplementary Fig. S4A). Adoptive transfer of WT B cells into B cell–deficient mice (B-cell KO–rescue group; Fig. 4A) showed a significant reduction in overall survival compared with the B-cell KO group, including compared with the WT group. B-cell immunophenotypic analysis revealed that adaptively transferred naïve B cells exhibited significantly higher expression of both inhibitory molecules CD155 and PD-L1 (Fig. 4B) in the tumor compared with those in the circulation (blood) and dCLNs. These data suggested that naïve B cells acquired a Breg phenotype within the tumor microenvironment. IL10-deficient B cells did not show any animal survival improvement (Supplementary Fig. S4B), suggesting that unlike other Breg populations (29), GBM-associated Bregs do not utilize IL10 as the main immunoregulatory mechanism.

GBM-associated Breg generation is an active process that requires MDSCs

Myeloid cells, especially monocytic MDSCs, are a prominent component of the GBM microenvironment (12, 30), and might play an important role in shaping the adaptive immune response to promote immune escape. We tested the ability of tumor cell–derived factors and MDSCs to promote Breg differentiation and function by culturing B cells with CT2A-conditioned media (CT2A-CM) or immunosuppressive MDSCs generated from BM cells using CT2A-CM (CT2A-AMDSCs; Supplementary Fig. S4C and S4D; Fig. 4C). Supernatants from CT2A-AMDSCs (CT2A-AMDSC-CM) were also tested. After 48 hours, B cells were collected and evaluated for their CD8+ T-cell–suppressive capability (Fig. 4C). Naïve B cells (Mock B) showed an enhancing effect in CD8+ T-cell activation (No B vs. Mock B, P < 0.005). B cells cocultured with MDSCs or MDSC-CM showed a significant suppressive effect compared with activated CD8+ T cells alone [No B vs. (CT2A-AMDSC)-B or (CT2A-AMDSC-CM)-B; P < 0.0001]. However, B cells incubated with CT2A-CM were not suppressive [No B vs. (CT2A-CM)-B, P 0.67] nor an activator of CD8+ T cells [Mock B vs. (CT2A-CM)-B, P > 0.005]. Because both CT2A-AMDSCs and CT2A-AMDSC-CM conferred regulatory function to B cells, we next evaluated whether this interaction required cell-to-cell contact. B-cell immunophenotype and function were analyzed in an MDSC/B-cell transwell system. We ensured that no contaminating MDSCs crossed the transwell 0.4-μm pore membrane (Supplementary Fig. S4E). We observed that the presence of PD-L1 on the surface of B cells was significantly increased after only 12 hours of culture, followed by increased expression of CD155, TGFβ (LAP), and IL10 at later time points (Fig. 4D; Supplementary Fig. S4F). These observations might suggest that the rapid acquisition of PD-L1 by B cells might mark the transition from naïve B cells to Bregs.

GBM-associated MDSCs induce Bregs via transfer of membrane-bound PD-L1

MDSCs can modulate the tumor microenvironment via cell-to-cell contact (i.e., PD-L1 and Fasl) by altering the surrounding microenvironment nutrient and metabolic profile (i.e., amino acid deprivation via arginase, ROS production) and by the production of immunosuppressive cytokines such as TGFβ or IL10 (31–33). In the context of cancer, MDSCs are known to secrete extracellular vesicles that can recapitulate many aspects of their immunosuppressive factors, mainly by transporting both membrane proteins and cytosolic factors to the target T cells (34–36). We observed that GBM MDSCs produced a significant amount of extracellular vesicles (Fig. 4E and F) able to cross the transwell 0.4-μm pore membrane. These extracellular vesicles were imaged by cryoelectronic microscopy (Fig. 4E), and the DLS analysis revealed the presence of vesicles with an average size of 188 nm (Fig. 4F). These MVs might correspond to exosomes (100–1,000 nm) rather than exosomes (40–100 nm; refs. 37, 38). To test whether naïve B cells could uptake the MDSC membrane–derived MVs, MDSCs were labeled with a lipophilic dye (CTV), and we repeated the transwell experiment. After 48 hours, B cells were harvested and evaluated for the acquisition of the CTV dye via flow cytometry. We observed that B cells acquired MDSC membrane–derived structures (CTV dye positive), which correlated with the presence of PD-L1 (Fig. 4G). Pretreatment of MDSCs with an inhibitor of MV release (GW4869) reduced CTV dye uptake by the B cells (Fig. 4G).

We next confirmed that B-cell uptake shown in Fig. 4G was due to MDSC-derived MVs because the majority of naïve B cells (>85%) directly incubated with MVs derived from MDSCs.
acquired the CTV signal after 12 hours, and >30% were also positive for PD-L1 (Fig. 4H). This suggested that the "rapid" increase of PD-L1 on B cells (Fig. 4D) was likely due to the uptake by the B-cell membrane of MDSC-derived MVs carrying PD-L1. MV uptake by B cells slightly increased expression levels of CD155 and TGFβ. No significant changes were observed for IL10 (Supplementary Fig. S4G). In CT2A-bearing mouse brains, monocytic MDSCs showed the highest PD-L1 expression in the tumor microenvironment (Fig. 4I), and MDSC-derived MVs carried PD-L1 on their membranes (not in the lumen; Fig. 4J). To verify this phenomenon, we knocked down PD-L1 in MDSCs using positively charged LNPCs to deliver siRNA. After 24 hours of transfection, MDSCs were harvested and tested for their PD-L1 transcripts using quantitative PCR (Supplementary Fig. S5A). MVs from transfected MDSCs (scramble siRNAs and pd-L1 siRNA) were harvested from supernatants and cultured with naive B cells for 12 hours. The presence of membrane PD-L1 on B cells was tested by flow cytometry. Our data showed that pd-L1 siRNA–transfected MDSC MVs transferred significantly reduced amounts of PD-L1 to B cells compared with the control scramble siRNA group (Fig. 5A). We confirmed that B cells cultured with MDSC-derived MVs suppressed CD8+ T-cell activation. However, blockade of PD-L1 on MVs before coculture with B cells significantly inhibited B-cell–suppressive effects on CD8+ T-cell expansion and effector phenotype (Fig. 5B). Altogether, these data suggested that MDSCs transferred PD-L1 to B cells via MVs, which conferred regulatory properties to B cells.

Brain-penetrant CSF-1R inhibitor (BLZ945) is known to reduce numbers of cancer-associated myeloid cells (39, 40). This diminished immune pressure on T cells and myeloid cells (41–43), and reduced PD-L1 expression by tumor-associated myeloid cells (ref. 39; Supplementary Fig. S5B). Our data showed that naive B cells transferred to B cell–deficient mice treated with BLZ945 (prior to B-cell injection) failed to acquire PD-L1 (Supplementary Fig. S5C). In support, in vivo blockade of PD-L1 upon radiotherapy (known to upregulate PD-L1 in myeloid cells; ref. 44) prevented the acquisition of regulatory properties by adoptively transferred B cells (Fig. 5C).

MV s can be acquired by B cells in an ATP-independent (4°C culture, passive fusion) or -dependent manner (37°C, active endocytosis; ref. 45). We observed that optimal MV uptake and PD-L1 transfer involved an ATP-dependent mechanism (37°C, Fig. 5D). An endocytosis inhibitor study showed that caveolae-mediated endocytosis could be a major mechanism of MV uptake by B cells. These results implied that PD-L1 on MVs was probably transferred to B cells via receptor-mediated endocytosis followed by endocytic recycling toward the plasma membrane of B cells (Fig. 5E; ref. 45).

Finally, the interaction of GBM-infiltrating Bregs with effector CD8+ T cells promoted overexpression of B-cell–derived TGFβ and IL10 (Supplementary Fig. S5D). This might reflect Bregs attempting to maintain the immunosuppressive environment by restraining CD8+ T-cell activation. Blockade of PD-L1 on tumor Bregs significantly reduced the expression of both TGFβ and IL10 by B cells. This observation highlights the magnitude of PD-L1 acquisition by B cells in controlling their own immune functions and the relevant role of MDSCs in inducing B cell–mediated immunosuppression.

Next, we aimed to elucidate whether human MDSCs and B cells could interact via transfer of MV-bound PD-L1. Human MDSCs were generated from peripheral blood monocytes using CM from patient-derived GBM xenograft lines (46): a classic subtype (GBM6), a proneural subtype (GBM12 and 43), and a mesenchymal subtype (MES83). We observed that MDSCs generated with CM from GBM12 and MES83 expressed high PD-L1 (Supplementary Fig. S6A). This observation correlated with acquisition of PD-L1 by B cells upon transwell coculture for 24 hours (Fig. 6A). B cells cultured with GBM12 and MES83-MDSCs were tested for their ability to suppress autologous CD8+ T cells in transwell experiments. Coculture with MES83-MDSCs provided the highest B-cell PD-L1 expression and significant immunosuppression (Fig. 6B and C). Because TAMs also express high PD-L1 and are also able to secrete MVs (47), we investigated whether CD163+ infiltrating macrophages from GBM patient tumors are able to transfer PD-L1 to the target cells. CD163+ infiltrating macrophages from freshly resected GBM tumors were isolated and tested for their ability to promote PD-L1+ Bregs. CD163+ macrophages labeled with the CTV dye (upper chamber) and PBMCs (lower chamber) were cocultured in a transwell system as in previous experiments. After 4 days, different immune cell populations were evaluated for the acquisition of the CTV dye and the presence of PD-L1. Foxp3+ Tregs and B cells showed the highest uptake compared with CD8+ T cells and myeloid cells (Fig. 6D; Supplementary Fig. S6B). CTV dye acquisition correlated with the presence of membrane PD-L1. CD33+ CD14+ myeloid cells expressed high PD-L1, but no sign of uptake (CTV signal) was observed (Supplementary Fig. S6B). Altogether, our observations suggest that GBM-associated MDSCs produced MVs that carry membrane-bound PD-L1, which are then taken up by target cells such as Bregs. Therefore, MDSCs may promote GBM by sharing their immunosuppressive factors with other cells prone to their reception.

**Discussion**

In the present study, we investigated the phenotype and function of GBM-infiltrating B cells. Whereas the role of other components of the tumor microenvironment such as MDSCs and Tregs is relatively well studied (4, 48–51), the involvement of B cells in GBM biology has not been thoroughly examined, limited only to the observation of B-cell infiltration in GBMs (14). Here, we demonstrated that B cells harvested from patient GBM tumors possessed a suppressive effect on activated CD8+ T cells, shown by the inhibition of CD8+ T-cell proliferation and further acquisition of an effector phenotype. Immunosuppressive functions of Bregs in cancer are also extended in the CD4+ T-cell compartment (52, 53). We observe a mild effect of glioma Bregs in suppressing activated CD4+ T cells. However, further studies should be pursued to deeply elucidate the role of glioma Bregs in CD4+ T-cell polarization and differentiation.

GBM-associated B cells showed a definite immunoregulatory phenotype as judged by expression of PD-L1, CD155, TGFβ, and IL10. These observations were also obtained in GL261 and C12A orthotopic models of GBM. This functional and immunophenotypic profile was exclusive to tumor-infiltrating B cells. We observed that local intratumoral depletion of B cells resulted in the improvement of animal survival. The therapeutic effectiveness was accompanied by increased intratumoral GzmB+ CD8+ T cells, suggesting that depletion of GBM-associated Bregs favored CD8+ T-cell activation and promoted their effector function. We did not observe significant animal survival benefit when B cells were depleted systemically. These results reinforce the hypothesis that
Figure 4.
MDSC converts naive B cells into Bregs via MVs. A, Survival of CT2A-bearing, B cell–deficient mice (B-cell KO) versus C57BL/6 (WT) mice. 5 × 10^6 naive B cells were adoptively transferred into B-cell KO mice (B-cell KO–rescue) and B-cell KO or WT groups (n = 10 mice/group). Survival curves were compared by log-rank test, and multiple comparisons were adjusted using the Bonferroni method. B, B-cell–deficient mice were injected with 5 × 10^6 B cells 7 days after tumor implantation, and B-cell phenotype was analyzed 3 days after. Percentage of PD-L1^+ and CD155^+ B cells in the brain, blood, or in the dCLNs was assessed in n = 4 mice/group. Differences were assessed by one-way ANOVA with Tukey multiple comparison test. C, MDSCs were generated using CT2A-CM for 6 days in the presence of GM-CSF (CT2A-MDSC). (Continued on the following page.)
B cells might present dual functions depending on their localization: protumorigenic functions in the tumor (Breg) versus antitumor functions in the periphery. Candolli and colleagues (54) reported similar observations when using systemic anti-CD20 depletion therapy in the GL261 GBM model. The authors observed that B cells played an important role in promoting antitumor responses by antigen presentation rather than Ab production. Our observations also contradict a computational analysis of GBM tumor-infiltrating immune cells, where the high magnitude of B cells predicted overall increased patient survival (55). Nevertheless, ex vivo tumor-infiltrating B cells from GBM tumor samples showed an immunosuppressive effect toward the CD8+ T-cell activation in agreement with previous observations (20, 21). Based on these data, we hypothesize that GBM and its microenvironment actively promote the conversion of B cells infiltrating the tumor into Bregs.

Several studies suggested that B-cell–deficient mice contribute to the understanding that the acquisition of the regulatory phenotype is an active process. In fact, adoptive transfer of naive B cells not only rescued the survival phenotype but also revealed that only tumor-infiltrating B cells showed high PD-L1 and CD155. These data suggest that a dynamic process might take place in the tumor microenvironment that converts naive B cells into GBM-associated Bregs. This hypothesis correlates with a current view of Breg differentiation, a biological mechanism that requires local immunologic activation (56). The GBM vicinity is an ideal microenvironment to attract and promote survival of potent immunoregulatory cells such as MDSCs or Tregs (12), and possibly Bregs. GBM cell membrane PD-L1 (own to release Bregs-inducing factors (20, 21). Other components of the tumor microenvironment such as myeloid cells promote activation of naive B cells and their differentiation toward an immunosuppressive functional profile. For instance, breast cancer–associated MDSCs have been reported to convert naïve B cells into immunosuppressive B cells able to inhibit T-cell responses (57). In GBMs, monotypic Ly6C-expressing MDSCs are the main component of the GBM microenvironment and represent approximately 40% of the total tumor mass (12, 13). We observed that B cells exhibited an immunoregulatory phenotype when cultured in the presence of GBM-associated MDSCs or MDSC supernatants, as they became capable of abrogating the CD8+ T-cell activation. The fast upregulation of MDSC-induced PD-L1 (unlike CD155, TGFβ, or IL10) on B cells after transwell culture with MDSCs led us to hypothesize that the initial increase of PD-L1 on B cells membrane might be partially due to a protein transfer from MDSCs. Chen and colleagues (58) report that tumor-derived exosomes, which are extracellular vesicles of endosalmon origin, can carry PD-L1 on their membrane surface. The authors observed that these "nanobodies" could suppress CD8+ T-cells via direct PD-L1/PD1 interaction remotely from the tumor site. This mechanism has been reported in GBM patients, where tumor-derived extracellular vesicles carrying PD-L1 on their surface can inhibit CD8+ T-cell activity (59). Accordingly, we observed that GBM-associated MDSCs secreted ~200 nm MVs carrying PD-L1 on their membrane. Our experiments showed that these MV shared cytoplasmic membrane as their cellular source, which are MDSCs. Naïve B cells were readily prone to uptake these shedding vesicles. We hypothesize that PD-L1 is carried by MDSC-derived MV membranes (because no detectable PD-L1 was observed in the MV lumen fraction) and is transferred to B cells. This hypothesis is supported by a recent report demonstrating that MDSC-derived extracellular vesicles can carry biologically relevant proteins (34, 35, 60). Accordingly, downregulation of PD-L1 expression on MDSCs significantly reduced PD-L1 presence on the surface of B cells and subsequently reduced Breg immunosuppressive effects on activated CD8+ T cells. This indicated that MDSC-derived MVs carrying PD-L1 are, at least in part, responsible for the presence of PD-L1 on B cells. This phenomenon might drive B cell–mediated immunosuppressive function in brain tumors.

Based on our results, one hallmark of GBM-associated Bregs is the overexpression of TGFβ and IL10 upon interaction with activated CD8+ T cells. This phenomenon might reflect an attempt of Bregs to maintain an immunosuppressive environment to prevent further CD8+ T-cell activation upon initial inhibitory contact. We are currently unable to determine whether MDSC-MV-bound PD-L1 transfer to B cells mediates TGFβ and IL10 upregulation, as it is unclear whether the PD-L1 downstream pathway is fully functional. Nevertheless, blockade of PD-L1 on tumor Bregs upon interaction with CD8+ T cells abrogated the overexpression of both TGFβ and IL10 by Bregs. One possible explanation might be the inability of CD8+ T cells to produce IFNγ upon contact with GBM Bregs. The lack of IFNγ might be the result of PD-L1-mediated inhibition (61) regulated by Bregs. Accordingly, PD-L1/PD-L1 interaction blockade promoted CD8+ T cell–derived IFNγ expression, which was associated with reduced expression of TGFβ and IL10 by B cells.

The mechanisms by which B cells infiltrate glioblastoma are unknown, and, albeit, Breg function might be relevant only in a...
reduced group of GBM patients. The overall study highlights a mechanism of MDSC cellular interaction with the adaptive immune system. The potential effect of GBM-associated myeloid cells in promoting Bregs via transfer of membrane-bound PDL1 might also occur in human myeloid cells. Both in vitro-generated MDSC-like cells (using MES83 mesenchymal cell line CM) and tumor-infiltrating CD163+ macrophages promoted increased PD-L1 in autologous B cells in transwell culture.
settings. CD163⁺ myeloid cell–derived MVs were preferentially up-taken by both B cells and Foxp3⁺ Tregs, suggesting the transfer of PD-L1–carrying MVs might represent a universal process of intercellular communication between regulatory cells.

In summary, we demonstrated that B cells infiltrating human and murine GBMs acquire the regulatory phenotype in the tumor milieu. This dynamic process was mediated, at least in part, by communication of MDSCs with B cells via MV-bound PD-L1 transfer from MDSCs. Our data suggest that Bregs play an
important role in GBM evasion from the surveillance of cytotoxic CD8$^+$ T cells, further highlighting the complexity of the immunosuppressive GBM network.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

Authors’ Contributions

Conception and design: C. Lee-Chang, A. Rashidi, J. Miska, I.V. Balyasnikova, M.S. Lesniak

Analysis and interpretation of data (e.g., statistical analysis, bioinformatics, computational analysis): C. Lee-Chang, A. Rashidi, J. Miska, P. Zhang, K.C. Pituch, D. Hou, M. Fischietti, M.S. Lesniak

Acquisition of data (provided animals, acquired and managed patients, provided facilities, etc.): C. Lee-Chang, A. Rashidi, J. Miska, P. Zhang, K.C. Pituch, D. Hou, C. Horbinski, A. Lopez-Rosas, Y. Han

Analysis and interpretation of data (e.g., statistical analysis, bioinformatics, computational analysis): C. Lee-Chang, J. Miska, P. Zhang, K.C. Pituch, T. Xiao, M. Fischietti, S.J. Kang, C.L. Appin, C. Horbinski, M.S. Lesniak

Writing, review, and/or revision of the manuscript: C. Lee-Chang, A. Rashidi, J. Miska, P. Zhang, K.C. Pituch, S.J. Kang, L.C. Platianis, I.V. Balyasnikova, M.S. Lesniak

Study supervision: C. Lee-Chang, L.C. Platianias, M.S. Lesniak

References


Acknowledgments

The authors would like to thank Katy McCartney and Rodrigo Javier at the Nervous System Tumor Bank (Feinberg School of Medicine, Northwestern University) for managing the interface between the clinic and the laboratory regarding GBM patients’ biological samples. Lastly, the authors are extremely grateful for the patients who consented to donate their biological samples. This study was supported by the following grants from the NIH (P50CA221747, R35CA197725, R01NS093903, and R01 NS087390); to M.S. Lesniak). C. Lee-Chang received support from a SPORE Career Enhancement Program (P50CA221747). The authors acknowledge support from the Structural Biology Facility at Northwestern University, the Robert H. Lurie Comprehensive Cancer Center of Northwestern University, and NCI CCSG P30 CA060553. The Gatan K2 direct electron detector was purchased with funds provided by the Chicago Biomedical Consortium with support from the Searle Funds at The Chicago Community Trust.

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Received April 1, 2019; revised July 23, 2019; accepted September 12, 2019; published first September 17, 2019.
Identification of Regulatory B Cells in GBM

Myeloid-Derived Suppressive Cells Promote B cell–Mediated Immunosuppression via Transfer of PD-L1 in Glioblastoma

Catalina Lee-Chang, Aida Rashidi, Jason Miska, et al.


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doi:10.1158/2326-6066.CIR-19-0240

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