## **JCI** insight

### The MIF promoter SNP rs755622 is associated with immune activation in glioblastoma

Tyler J. Alban, ..., Manmeet S. Ahluwalia, Justin D. Lathia

JCI Insight. 2023;8(13):e160024. https://doi.org/10.1172/jci.insight.160024.

Research Article Immunology Oncology

Intratumoral heterogeneity is a defining hallmark of glioblastoma, driving drug resistance and ultimately recurrence. Many somatic drivers of microenvironmental change have been shown to affect this heterogeneity and, ultimately, the treatment response. However, little is known about how germline mutations affect the tumoral microenvironment. Here, we find that the single-nucleotide polymorphism (SNP) rs755622 in the promoter of the cytokine macrophage migration inhibitory factor (MIF) is associated with increased leukocyte infiltration in glioblastoma. Furthermore, we identified an association between rs755622 and lactotransferrin expression, which could also be used as a biomarker for immune-infiltrated tumors. These findings demonstrate that a germline SNP in the promoter region of MIF may affect the immune microenvironment and further reveal a link between lactotransferrin and immune activation.







# The MIF promoter SNP rs755622 is associated with immune activation in glioblastoma

Tyler J. Alban, <sup>1,2</sup> Matthew M. Grabowski, <sup>3</sup> Balint Otvos, <sup>3</sup> Defne Bayik, <sup>1</sup> Wesley Wang, <sup>4</sup> Ajay Zalavadia, <sup>1</sup> Vlad Makarov, <sup>2</sup> Katie Troike, <sup>1</sup> Mary McGraw, <sup>3</sup> Anja Rabljenovic, <sup>1</sup> Adam Lauko, <sup>1</sup> Chase Neumann, <sup>1</sup> Gustavo Roversi, <sup>1</sup> Kristin A. Waite, <sup>5</sup> Gino Cioffi, <sup>5</sup> Nirav Patil, <sup>6</sup> Thuy T. Tran, <sup>7</sup> Kathleen McCortney, <sup>8</sup> Alicia Steffens, <sup>8</sup> C. Marcela Diaz, <sup>2</sup> J. Mark Brown, <sup>1,9,10</sup> Kathleen M. Egan, <sup>8</sup> Craig M. Horbinski, <sup>8</sup> Jill S. Barnholtz-Sloan, <sup>5</sup> Prajwal Rajappa, <sup>4</sup> Michael A. Vogelbaum, <sup>11</sup> Richard Bucala, <sup>7</sup> Timothy A. Chan, <sup>2,3,9,10</sup> Manmeet S. Ahluwalia, <sup>12</sup> and Justin D. Lathia<sup>1,3,9,10</sup>

'Department of Cardiovascular & Metabolic Sciences and Imaging Core, Lerner Research Institute, <sup>2</sup>Center for Immunotherapy and Precision Oncology, and <sup>3</sup>Rose Ella Burkhardt Brain Tumor and Neuro-Oncology Center, Cleveland Clinic, Cleveland, Ohio, USA. <sup>4</sup>Nationwide Children's Hospital, Institute for Genomic Medicine, Departments of Pediatrics and Neurological Surgery, The Ohio State University Wexner Medical Center, Columbus, Ohio, USA. <sup>5</sup>Division of Cancer Epidemiology and Genetics, Trans-Divisional Research Program, Center for Biomedical Informatics and Information Technology, National Cancer Institute, Bethesda, Maryland, USA. <sup>6</sup>University Hospitals Research and Education Institute, Cleveland, Ohio, USA. <sup>7</sup>Yale School of Medicine and Yale Cancer Center, New Haven, Connecticut, USA. <sup>8</sup>Departments of Pathology and Neurosurgery, Northwestern University, Feinberg School of Medicine, Chicago, Illinois, USA. <sup>9</sup>Cleveland Clinic Lerner College of Medicine at Case Western Reserve University, Cleveland, Ohio, USA. <sup>10</sup>Case Comprehensive Cancer Center, Cleveland, Ohio, USA. <sup>11</sup>Departments of Cancer Epidemiology and Neuro-Oncology, H. Lee Moffitt Cancer Center, Tampa, Florida, USA. <sup>12</sup>Miami Cancer Institute, Baptist Health South Florida, Miami, Florida, USA.

Intratumoral heterogeneity is a defining hallmark of glioblastoma, driving drug resistance and ultimately recurrence. Many somatic drivers of microenvironmental change have been shown to affect this heterogeneity and, ultimately, the treatment response. However, little is known about how germline mutations affect the tumoral microenvironment. Here, we find that the single-nucleotide polymorphism (SNP) rs755622 in the promoter of the cytokine macrophage migration inhibitory factor (*MIF*) is associated with increased leukocyte infiltration in glioblastoma. Furthermore, we identified an association between rs755622 and lactotransferrin expression, which could also be used as a biomarker for immune-infiltrated tumors. These findings demonstrate that a germline SNP in the promoter region of *MIF* may affect the immune microenvironment and further reveal a link between lactotransferrin and immune activation.

Conflict of interest: MSA received grants/research support from AstraZeneca, BMS, Bayer, Incyte, Pharmacyclics, Novocure, Mimivax, and Merck. MSA received honoraria or consultation fees from Bayer, Novocure, Kiyatec, Insightec, GSK, Xoft, Nuvation, Cellularity, SDP Oncology, Apollomics, Prelude, Janssen, Voyager therapeutics, Caris Lifesciences, Annheart Therapeutics, Viewray, Forma Therapeutics, Astrazeneca, Karyopharm, BMS, Tocagen, Abbvie, VBI Vaccines, Flatiron, Bayer, and Varian Medical Systems. MSA is a stock shareholder with Doctible, Mimivax, Cytodyn, MedInnovate Advisors LLC.

**Copyright:** © 2023, Alban et al. This is an open access article published under the terms of the Creative Commons Attribution 4.0 International License.

**Submitted:** March 9, 2022 **Accepted:** May 25, 2023 **Published:** May 30, 2023

Reference information: /Cl Insight. 2023;8(13):e160024. https://doi.org/10.1172/jci. insight.160024.

#### Introduction

Well-known somatic drivers of glioblastoma (GBM) heterogeneity, including *PDGFRA*, *IDH1*, *EGFR*, and *NF1*, have altered how we treat patients, diagnose disease, and design clinical trials (1, 2). While many groups have sought to identify germline variants associated with disease prevalence and outcomes, it is still unclear how germline variants alter the tumoral heterogeneity and immune microenvironment. Here we focus on understanding the tumoral effects of 1 well-characterized germline single-nucleotide polymorphism (SNP), rs755622, which has been shown to be associated with increased expression of the cytokine macrophage migration inhibitory factor (MIF) (3–20). MIF is an important microenvironmental factor in glioma, and we previously identified an immune-suppressive pathway in GBM that is driven by secreted MIF from cancer stem cells (CSCs) that in turn activates myeloid-derived suppressor cells (MDSCs) (21). Recent work from our laboratory and others has shown that MDSCs are increased in the circulation and tumor microenvironment (22), they portend a poor prognosis (23), their expansion can be driven by CSC-derived MIF (21, 23), and they can be reduced by MIF neutralization (either genetically or pharmacologically) (24, 25). Furthermore, *MIF* has been studied in a variety of cancers in the context of inflammation and has been found to regulate immune activity (3–20).

1



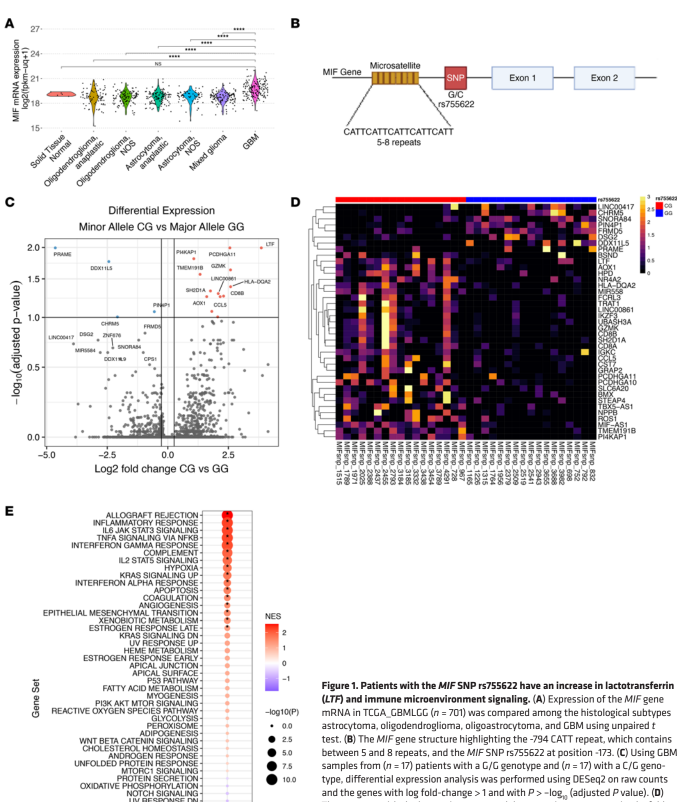
MIF expression is influenced not only by disease state but also by functional germline genetic polymorphisms. The most notable is a MIF promoter SNP, rs755622, which has been associated with multiple inflammatory conditions such as lupus, rheumatoid arthritis, septic shock, and cardiovascular disease (3, 5, 8, 13, 15). This MIF SNP is in linkage disequilibrium with the presence of 7 CATT repeats at the –794 promoter microsatellite, which leads to tighter binding by the transcription factor ICBP90 when compared with the more commonly occurring 5 or 6 CATT repeats (26). Accordingly, rs755622 is commonly analyzed in place of the –794 CATT microsatellite (15). While the minor allele frequency of the rs755622 MIF SNP ranges from 15% to 20% in White individuals and > 45% in individuals of African descent, it has not been associated with GBM growth or survival in large-scale genome-wide association studies (27). Given the role of MIF in immune response, we hypothesized that this MIF SNP may be associated with the immune microenvironment of GBM. Here, we determined that patients with the MIF SNP rs755622 have an altered tumor microenvironment characterized by increased lymphocyte infiltration and enhanced lactotransferrin (LTF) expression. Furthermore, high LTF expression can serve as an indicator for a subset of tumors that have high lymphocyte infiltration.

#### Results

Patients with the MIF SNP rs755622 have an increase in LTF expression and increased immune microenvironment signaling. Based on our previous assessments of MIF as a driver of CSC/MDSC-mediated communication (21), we assessed overall MIF expression levels across brain tumors and found elevated MIF in isocitrate dehydrogenase (IDH) WT GBM patient tumor samples when compared with patients with lower-grade (IDH mutant astrocytomas and oligodendrogliomas) gliomas and normal tissue using TCGA data (Figure 1A). Based on these information, we hypothesized that patients with GBM may have an increased prevalence of the regulatory MIF rs755622 nucleotide -173 G/C SNP (Figure 1B), which has been shown to be 1 factor with the ability to increase MIF expression. We assessed the rs755622 MIF SNP across 3 separate, annotated clinical cohorts of patients with GBM (total of 966 patients, including 449 from Cleveland Clinic, 386 from Moffitt Cancer Center, and 131 from Case Western Reserve University/University Hospitals of Cleveland). Our analysis of individual and combined cohort statistics revealed significant differences in the frequency of key prognostic indicators among cohorts (Karnofsky Performance Score [KPS], total surgical resection, receipt of standard of care [SOC], and recurrence) across the 3 cohorts (Supplemental Figure 1, A and B; supplemental material available online with this article; https://doi.org/10.1172/jci.insight.160024DS1). Across all cohorts, we genotyped patients via PCR and found a similar frequency of MIF SNP rs755622 major (G/G) and minor (C/C or C/G, noted as C/\*) allele-containing genotypes in each (Supplemental Figure 1C). Furthermore, these frequencies matched the expected frequencies reported in the 1000 Genomes Project in people of European ancestry (Supplemental Figure 1C). Previous reports (28) have also shown a strong linkage disequilibrium between the rs755622 SNP and the CATT repeats rs5844572, so we further assessed the CATT repeat length in patients with available DNA (Supplemental Figure 2, A and B). In this analysis, we found a > 90% cosegregation between CATT repeats 7-8 (rs5844572) and the MIF SNP (rs755622), in alignment with previous studies (28). When evaluated according to major clinical prognostic indicators for GBM, we observed significant associations of sex and receipt of SOC with the rs755622 MIF SNP genotype using descriptive analysis (Supplemental Figure 1, B, D, and E, and Supplemental Figure 3). Univariate and multivariable survival analyses did not demonstrate any differences in overall or progression-free survival (PFS) according to rs755622 MIF SNP genotype in any individual cohort or when data from all 3 cohorts were combined (Supplemental Figure 1, D-F, and Supplemental Figures 3 and 4).

When analyzing individual cohorts by univariate analyses and combined analyses, we observed no significant difference in GBM incidence or patient survival between the rs755622 MIF SNP genotypes; however, we hypothesized that there may be differences in tumor and microenvironment interactions between genotypes given the association of the rs755622 MIF SNP with inflammation in nononcologic conditions (Supplemental Table 1). To explore this possibility, we performed RNA-Seq on 17 patients with primary, untreated GBM from each MIF genotype in our Cleveland Clinic cohort who had similar clinical parameters and outcomes (Supplemental Table 2). In selecting this cohort, we included patients with good prognosis (greater than 6 months PFS from diagnosis) and poor prognosis (less than 6 months PFS from diagnosis). We balanced our data set in this way to ensure that we did not bias our results toward an immunologic phenotype based on observations that patients with better outcomes have enriched lymphocytes (22, 29, 30). Differential gene expression analysis revealed that the rs755622 minor allele patients (e.g., -173 C SNP) had an enrichment in immune cell–related genes, including LTF, GZMK, HLA-DQA2, CD8B, and CCL5 (Figure 1, C and D).





7.5 10.0

between 5 and 8 repeats, and the MIF SNP rs755622 at position -173. (C) Using GBM samples from (n = 17) patients with a G/G genotype and (n = 17) with a C/G genotype, differential expression analysis was performed using DESeq2 on raw counts and the genes with log fold-change > 1 and with  $P > -\log_{10}$  (adjusted P value). (D) The 25 genes with the largest increases and decreases in gene expression by fold change are shown via heatmap (rs755622; red, CG; blue, GG; ID, colors indicate individual samples with column being a unique sample). Heatmap rows are clustered using hierarchical clustering; transcripts per million are scaled to rows for heatmap color scale. (E) GSEA was performed using Hallmark gene sets on the differentially expressed gene list comparing the C/G genotype to the G/G genotype sorted by log fold-change for gene-rank position. Normalized enrichment scores (NES) are shown in red/blue, with red enriched in C/G and blue enriched in G/G, while the size of the circle represents the  $-\log_{10} P$  value as determined by GSEA.

CG vs. GG

**GSEA Enrichment Score** 

(\* = P < 0.05)

UV RESPONSE DN

MYC TARGETS V1 SPERMATOGENESIS MITOTIC SPINDLE G2M CHECKPOINT E2F TARGETS

MYC TARGETS

DNA HEPAIH
TGF BETA SIGNALING
HEDGEHOG SIGNALING
PANCREAS BETA CELLS
BILE ACID METABOLISM



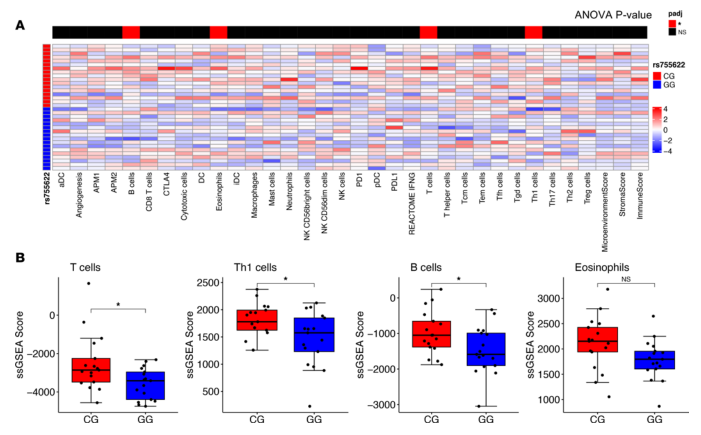
Gene-set enrichment analysis (GSEA) utilizing Hallmark curated gene sets identified a significant increase in inflammatory pathways, including Interferon Gamma Response and Allograft Rejection, in *MIF* SNP minor allele–containing tumors (Figure 1E). Additionally, ingenuity pathway analysis (IPA) showed similar findings, with increased innate immune response and prostaglandin signaling among the top enriched pathways of patients with the *MIF* SNP minor allele (Supplemental Figure 5). In seeking to better understand the individual immune cell types that were different between genotypes, we used single-sample GSEA (ssGSEA) deconvolution with curated gene sets related to immune cells and immunotherapy. In these analyses, we observed a significant increase in cell signatures associated with an immune response in patients with the minor allele compared with those with the major allele; significance was calculated by 2-way ANOVA and is indicated by the *P* value bar above each column of the heatmap (Figure 2, A and B).

Of the differentially expressed genes, *LTF* was the most significantly upregulated in patients with the minor allele and positively correlated with increased T cell and immune responses (Figure 1C). LTF is considered a key factor in first-line immune defense against bacteria, yeast, viruses, fungi, and parasites and may additionally contribute to antitumor response (31–33). LTF is an acute-phase immune mediator released from neutrophils and participates in the switch from innate to adaptive immune response. LTF signals through TLRs in myeloid cells to activate NF-κB and CD40 expression and to promote the initiation of antitumor immune responses (34, 35). It has also been associated with M1 polarization and, in a pancancer study, negatively correlated with tumor mutational burden (36). Taken together, these analyses suggest that, while the rs755622 *MIF* SNP did not associate with differences in GBM incidence or survival, rs755622 did correlate with a difference in the immune cell composition of the tumor microenvironment.

Immunofluorescence staining confirms enhanced T cell infiltration and CD8<sup>+</sup> T cell activation in patients with GBM with the MIF SNP rs755622. To further interrogate the immune cell differences between patients with differing MIF genotypes, we utilized matched tissue from 22 of the patients from our RNA-Seq cohort (12 minor allele C/\* and 10 major allele G/G). Staining for LTF confirmed the RNA-Seq analysis and showed increased LTF protein in minor allele patients (Figure 3, A–C). The percentage of LTF-expressing cells was further enhanced in minor allele patients with shorter survival spans, based on an overall survival of less than the median of the 34 samples in this cohort (Figure 3D). Additional assessment of T cell populations identified significant increases in total T cells in minor allele patients (Figure 3, E and F), and this increase appeared to mostly correspond to CD8<sup>+</sup> T cells, based on similar results from quantifying total CD3<sup>+</sup> or CD8<sup>+</sup> cells (Figure 3, F and G). While analysis of the activation marker CD107a on CD8<sup>+</sup> T cells showed no difference between patients with different prognoses (Figure 3H), CD107a was significantly increased in CD8<sup>+</sup> T cells in patients with the minor allele (Figure 3I). These data support the conclusion that immune infiltration by a cytotoxic T cell population and LTF expression are increased in the tumor microenvironment of patients with the minor allele.

While we observed differences in the lymphoid compartment of major and minor allele patients by RNA-Seq and validated these in patient samples, these initial assessments did not focus on specific myeloid cell subtypes that are known to be involved in GBM immune suppression (including microglia, monocytes, macrophages, and MDSCs). Again using matched samples from the patients included in the RNA-Seq study, we next stained for CD4<sup>+</sup> T cells and these myeloid cell subtypes (n = 12 C/\* and n = 10 G/G). First, we identified individual cell subtypes using the top quartile for lineage-specific markers (CD4+ T cells: CD4+CD11b-; macrophages: CD11b+CD68+HLA-DR+; microglia: P2RY12+; MDSCs: CD11b+C-D74\*CD68-HLA-DR-; monocytes: CD11b\*HLA-DRA\*P2RY12-CD68-) (Figure 4A). Heatmap representations of cell type markers are shown for all cells in the data set using the average cell MFI for each marker (Figure 4A). Representative images for each marker are shown, highlighting specific cell types from 1 sample section containing all cell types to demonstrate staining specificity for markers for the different cell types (Supplemental Figure 6A). Quantification revealed decreased macrophages in minor allele patients but no other major changes in MDSCs or the ratio of MDSCs to CD8<sup>+</sup> T cells, microglia, monocytes, or CD4<sup>+</sup> T cells between genotypes (Figure 4B). Due to the quantitative difficulties of assessing all myeloid cell lineages in an individual panel with lymphoid populations, we correlated cell types across samples per genotype (Figure 4, C and D). Using this approach, we found striking differences between genotypes, with the minor allele patients exhibiting an increase in CD8<sup>+</sup> T cells and reductions in the myeloid compartment. Additionally, LTF positively correlated with microglia and negatively correlated with the ratio of MDSCs to CD8<sup>+</sup> T cells. These data lead to the overall conclusion that minor allele patients have increased lymphocyte infiltration with reduced macrophage content and that LTF is associated with increased CD8+ T cells.





**Figure 2. Patients with the MIF SNP rs755622 have an increased immune infiltration.** (**A**) ssGSEA analysis was performed using the R package GSVA for cell type gene signatures, while xCell was utilized to estimate the overall immune, microenvironment, and stromal scores. ANOVA was performed to compare the deconvolution scores between patients with C/G and G/G genotypes, with the *P* value shown above the heatmap of the scaled scores. (**B**) ssGSEA scores for significant populations are shown as a box plot with unpaired *t* test *P* values shown for each.

In an effort to better understand how the immune microenvironment plays a role in treatment response, we next developed a cohort of n = 8 recurrent glioma samples for spatial profiling using nanostring spatial genomics platform. In this study, we focused on areas confirmed to have abnormal nuclear cellularity, suggesting cancer cells were targeted and is responsive to treatment. We then compared differential expression of these treatment responsive regions versus progressive regions and found that immune-related genes, including LTF, were increased in treatment reactive regions (Figure 4E). Ultimately, cases that presented clinically as treatment reactive also had increased enrichment for LTF expression (Figure 4F).

Patients with GBM with high LTF expression are immunologically activated. Seeking to expand on these initial observations, we sought to identify the rs755622 SNP in The Cancer Genome Atlas (TCGA) data set. However, we were not able to detect the SNP in this data set because the marker is too far upstream to have read coverage in whole-exome sequencing data. Given the strong correlation between the MIF minor allele and LTF expression, we used LTF expression as a surrogate of MIF genotype and interrogated immune changes in the context of LTF expression. Notably, we found a similar differential expression profile between LTF-high (top 25%) and LTF-low (bottom 25%) patients in the TCGA compared with what we found between genotypes in our data set (Figure 5A). In agreement with our initial assessment, GSEA revealed an increase in immune activation pathways in the LTF-high patients, including Allograft Rejection and Complement Signaling, and a reduction in cancer-related pathways, including Mitotic Spindle and Myc Targets (Figure 5, B and C). To further identify cell type estimates between LTF-high and LTF-low samples, we performed deconvolution analyses and found increased immune and microenvironment scores, IFN-γ score, and macrophage content (Figure 6, A and B). However, we did not see significant differences between LTF-high and LTF-low samples in many T cell subsets, such as T central memory cells, T effector memory cells, T follicular helper cells, γδ T cells, and Th1 cells.

rs2096525 serves as a surrogate for rs755622 to identify increased immune activation in GBM. To determine whether these findings with respect to LTF gene signature recapitulate the rs755622 genotype, we examined possible associations with the rs2096525 SNP, which is in linkage disequilibrium with rs755622 but located



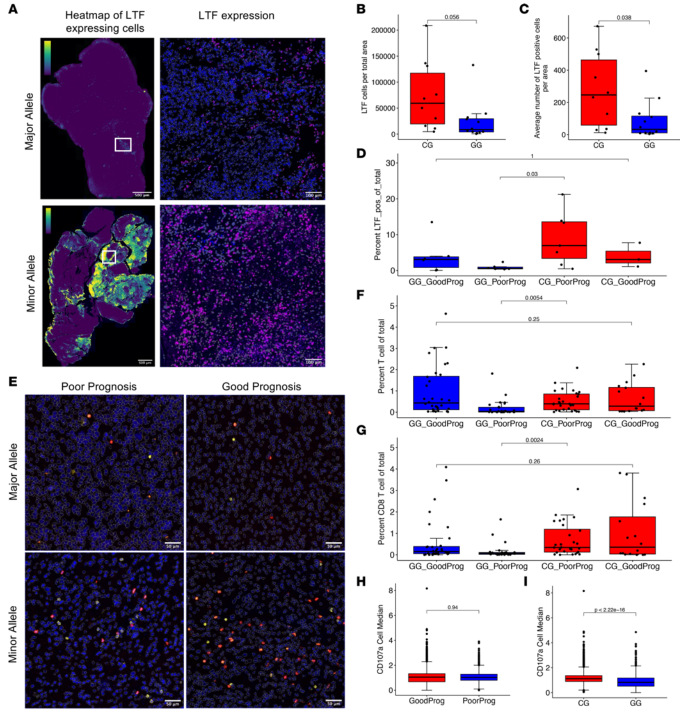
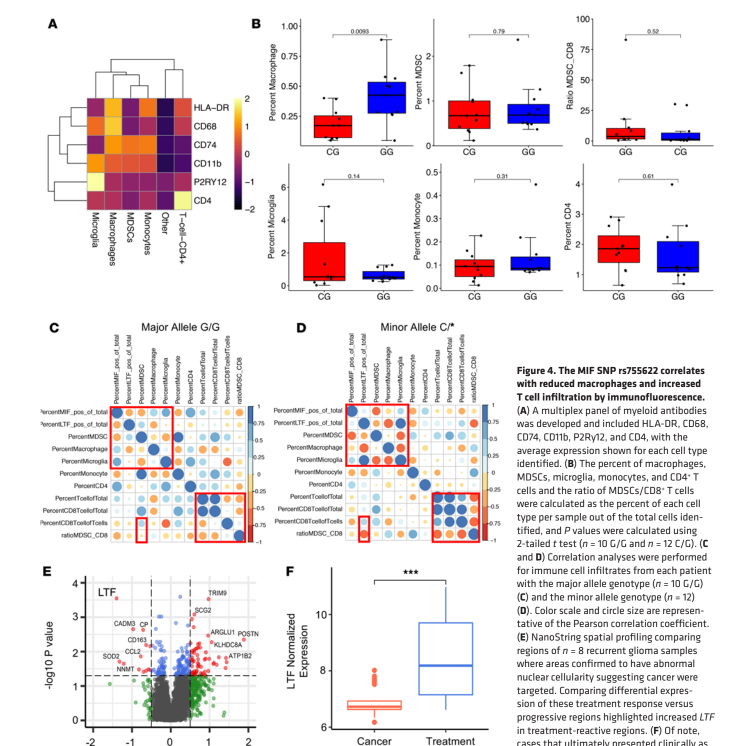


Figure 3. Immunofluorescence confirms enhanced T cell infiltration and CD8 $^{\circ}$  T cell activation in patients with GBM with the *MIF* SNP rs755622. (A) *LTF* expression was analyzed by immunofluorescence of matched tissue for n =22 samples from the RNA-Seq analysis to compare patients with the minor allele (C/G) (n = 10) and those with the major allele (G/G) (n = 12), with representative images of the data set shown as a whole view of the slide in the heatmap images (left column: yellow, increased density of staining). In a 20 $^{\circ}$  image of the slide, cells are outlined in white, with nuclei marked by DAPI and LTF pseudocolored purple. Left column: Full slide scans, zoomed to see the entire section. Scale bar: 500 μM. LTF is pseudocolored from purple to yellow scaled from 0 to 0.2. Right column: Representative zoomed in regions of major allele (top) and minor allele (bottom), where nuclei marked with blue DAPI and LTF colored purple. Scale bar: 100 μM. (**B**) The average LTF expression (mean fluorescence intensity) of cells per sample was compared between each genotype and the negative secondary-only control (n = 10 C/G and n = 12 G/G). (**C**) The quantity of LTF $^{\circ}$  cells per total area was measured and then compared between the C/G and G/G genotypes using unpaired t test (n = 10 C/G and n = 12 G/G). (**D**) The percent of LTF $^{\circ}$  cells of the total cells per sample was compared between each genotype group and further subdivided into prognostic categories based on greater than or less than median overall survival (poor prognosis contains n = 6 G/G and n = 7 C/G, good prognosis contains n = 6 G/G and n = 3 C/G). (**E**) Staining for CD3 and CD8 markers to determine CD8 $^{\circ}$  T cell infiltration, with representative images of each genotype and prognosis group (CD3, yellow; CD8, red; DAPI, blue). Scale bar: 100 μM. (**F** and **G**) Percent of T cells (**F**) and CD8 $^{\circ}$  T cells (**G**) of total cells per sample comparing the genotype/prognosis categories only. (**H**) CD107a expression of CD8 $^$ 





Progression

within the first or second MIF intron. We interrogated the TCGA GBM whole-exome sequencing data set for the rs2096525 SNP and found the minor allele to be present in approximately 12% of patients with GBM (Supplemental Figure 7A). Differential expression analysis did not identify LTF, MIF, or the MIF receptors (Supplemental Figure 7B); however, functional GSEA did identify a similar increase in immune activation (Supplemental Figure 7, C and D). Additionally, a deconvolution analysis demonstrated similar increases in macrophages, neutrophils, and T cells for this SNP as observed by the rs755622 RNA-Seq study of Cleveland Clinic Foundation patients (Supplemental Figure 7E).

Reaction

-2

Treatment

Reaction

-1

0

Log2

Fold Change

2

Cancer

Progression

cases that ultimately presented clinically as

for LTF (\*\*\*P < 0.001, n = 8).

treatment reactive had increased enrichment



#### Discussion

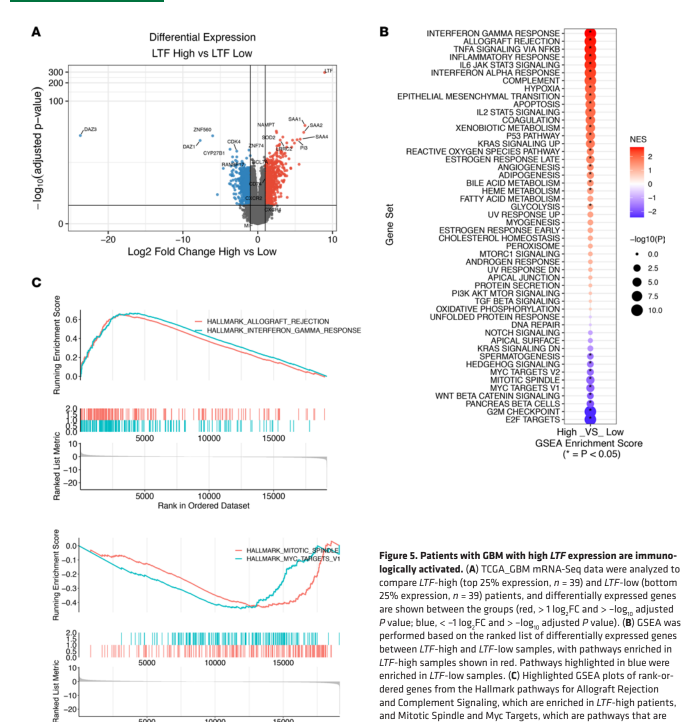
MIF, considered the first active cytokine discovered (37), has been extensively studied in the context of immune activation and the inflammatory response, as well as in tumor biology, where it has been shown to drive cancer cell proliferation and the generation of a tumor-promoting immune microenvironment (38). In GBM, functions of MIF include enhancement of CSC maintenance (39, 40), resistance to therapies including SOC chemotherapy temozolomide (41) and the antiangiogenic agent bevacizumab (42), and alteration of growth factor receptor signaling (34). However, the potential functional consequences of common MIF promoter polymorphisms, including the -173 SNP (rs755622), have not been well studied in the context of the inflammation associated with malignancies. Furthermore, the effects of MIF promoter polymorphisms have been studied even less often in CNS-associated disorders due to the strong expression quantitative trait locus (eOTL) in CNS tissues; however, we know that the immune cells play an integral role in CNS tumor progression and that they are in eQTL (Supplemental Figure 8). While the rs755622 SNP is associated with numerous inflammatory conditions and certain cancers, particularly those sensitive to immunotherapies (7, 38, 43, 44), we found no correlation with GBM incidence or prognosis in response to SOC therapy across 3 studied cross-institutional cohorts. This finding extended to the functional MIF promoter -794 CATT microsatellite (rs5844572) that is in linkage disequilibrium with the rs755622 SNP. We also observed no major difference in MIF gene expression level between genotypes, likely due to the elevated level of MIF in GBM or by the treatment of many patients with GBM with dexamethasone, which increases MIF production (45). However, we found evidence for distinct tumor immune microenvironments between SNP genotypes, with an increase in CD8+T cells in the minor allele patients. This enhancement in immune response parameters correlated with an enhancement in LTF expression in the minor allele patients.

Our initial assessment of *MIF* genotypes revealed an association between the minor SNP allele and *LTF* that has not been previously described. In nonpathophysiological conditions, LTF is an iron-binding glycoprotein that functions to protect against pathogens and has been shown to have antiin-flammatory activity. In cancer, LTF has been described to function in an antiproliferative manner. *LTF* expression is reduced in GBM compared with lower-grade brain tumors (46) and can inhibit GBM cell proliferation (47). LTF has also been studied as a nanoparticle carrier for a variety of preclinical cancer therapies, including in GBM, where it is able to penetrate the brain (48). In the context of MDSCs, we found that MIF enhances MDSC function in GBM and that LTF can induce MDSCs in pathological neonatal inflammatory conditions (49). While the MIF and LTF pathways have not been directly linked, our data suggest a likely interaction. *LTF* and *MIF* are likely not coregulated due to their location on separate chromosomes and their distinct transcription factors, but future studies interrogating the molecular relationship between LTF and MIF may provide additional insight into signaling networks that functionally link these 2 proteins in GBM.

The association of *LTF* with higher microenvironmental immune response and improved prognosis was further highlighted by our spatial profiling analysis. While both groups contained areas of cancer cellularity, treatment-reactive processes such as pseudoprogression have been associated with abundant global immune infiltration (50). Furthermore, patients experiencing reactions to treatment have better survival outcomes (51–53). Based on these results, it's reasonable to hypothesize that *LTF* expression could be prognostic in the immunotherapeutic setting. Together, these data indicate a distinct activated immune environment in patients containing the minor allele of the *MIF* SNP that is associated with *LTF* expression and CD8<sup>+</sup> T cell infiltration.

Numerous groups have reported on the association between CD8<sup>+</sup> T cell infiltration and prognosis in GBM, with varying results; however, a majority of studies appear to show a positive association between T cell infiltration and overall survival (54–56). In more recent literature, a more nuanced point of view has emerged recognizing the specific impact of proliferating CD8<sup>+</sup> T cells and/or the ratio of CD8<sup>+</sup>/CD4<sup>+</sup> T cells on survival in patients with GBM (57, 58), supporting a necessary balance within the cellular immune response in GBM to improve current treatment strategies. The challenge of predicting outcomes with simple cell enumeration in a complex disease such as GBM is reflected in our study, as the *MIF* SNP was associated with increased CD8<sup>+</sup> T cells, but not outcome, in our patient population. However, tumor-infiltrating lymphocytes (TILs), and CD8<sup>+</sup> TILs in particular, are known to be critical for limiting tumor progression (57). In the context of immunotherapy, microenvironments high in CD8<sup>+</sup> TILs are typically referred to as "hot" tumors and have been demonstrated to have a higher response rate to immunotherapies. While this phenotype is only seen in approximately 15% of GBM cases, this observation could indicate an immunotherapy-responsive subtype of GBM, which future studies should explore.





In analyzing the MIF SNP rs2096525, which is in linkage disequilibrium with rs755622, we were initially surprised to not identify overlap of top differentially expressed genes with our rs755622 RNA-Seq cohort. However, it is important to note that we utilized TruSeq methods of library preparation along with a deep-sequencing approach. Additionally, TCGA-GBM samples have many batch effects such as institution, sequencers, and potential necrotic samples that are difficult to control for in a comparison such as this. Together, these factors make it challenging to interpret the direct findings with this SNP from TCGA. However, the use of broader approaches such as GSEA did produce overlapping signatures similar to that of our cohort. This may indicate that these samples are overall more immune activated, but mechanistic studies of LTF expression in relation to the MIF SNP rs755622 and MIF itself are needed to understand whether LTF plays a direct role in immune activation.

enriched in LTF-low patients.

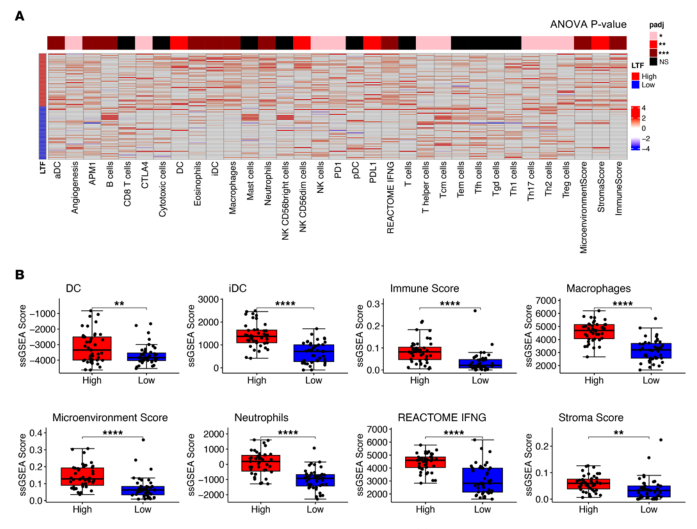
and Mitotic Spindle and Myc Targets, which are pathways that are

10000

Rank in Ordered Dataset

15000





**Figure 6. Patients with GBM with high** *LTF* **expression have increased immune infiltration.** (**A**) Deconvolution analysis of the samples belonging the *LTF*-high (n = 39) and *LTF*-low (n = 39) groups showed increased immune cell type infiltration and increased immune scores, with ANOVA for multiple comparisons showing the *P* **value** by heatmap coloring. (**B**) Individual plots for significantly different estimated cell types and scores shown as individual box plots with unpaired *t* test (n = 39 samples per group).

Utilizing both RNA-Seq and matched tissue samples, we identified that MIF SNP minor allele patients with increased *LTF* expression also had an increase in CD8<sup>+</sup> T cells and a reduction in macrophages, with no change in MDSCs. We also did not observe a consistent change in tumor-associated macrophages, and this could be due to a limitation of deconvolution methods in distinguishing myeloid subtypes. Taken together, this immune microenvironment may be more conducive to immune-activating strategies, and this should be assessed in future clinical trials. While our initial analysis revealed a high correlation between the *MIF* minor allele and elevated *LTF*, the association between the *MIF* SNP minor allele and *LTF* is indirect. This was due to the inability to efficiently identify *MIF* SNP status in large genomic data sets based on its location in the promoter region, which is not covered by whole-exome sequencing. Nonetheless, utilizing *LTF* at a median cut-off did not yield the same results as the top quartile of *LTF*, which more closely represents the frequency of the *MIF* SNP minor allele. Another limitation of our findings is that the *MIF* SNP has not been functionally characterized but is in linkage disequilibrium with the *MIF* –794 CATT repeat, which is associated with an increase in *MIF* production in immune cells and brain tissues.

The known genetic determinants of immunotherapy response in gliomas, including somatic mutations in *IDH*, are limited, and the present findings identify a common germline SNP linked to potential immunologic differences that may help inform clinical decision-making and be leveraged for the development of more effective immunotherapies.



#### **Methods**

DNA isolation and quantification. Genomic DNA was extracted from the peripheral blood of patients with GBM using a Qiagen DNeasy Blood & Tissue Kit following the manufacturer's protocols. DNA purity and concentration were measured using a Thermo Fisher Scientific NanoDrop spectrophotometer.

SNP genotyping. MIF SNP genotyping was performed using PCR amplification and subsequent restriction enzyme digestion with AluI. PCR was performed using Accuprime Pfx DNA polymerase (Thermo Fisher Scientific, catalog 12344-024) using 0.2  $\mu$ mol forward primer (5'-CCCAAAGACAGGAGGTAC-3') and 0.2  $\mu$ mol reverse primer (5'-ATGATGGCAGAAGGACCAG-3'). PCR was run as follows: 95°C for 5 minutes, followed by 35 cycles of 95°C for 30 seconds, 60°C for 45 seconds, and 68°C for 1 minute. Following the 35 cycles, there was a final 68°C elongation step for 5 minutes, followed by storage at 4°C. After amplification, the PCR product was confirmed on a 1% agarose gel by identification of an approximately 500 bp product. After confirmation, 10  $\mu$ L of PCR product was mixed with 2  $\mu$ L 10× CutSmart buffer, 1  $\mu$ L AluI, and 7  $\mu$ L water and digested at 37°C for 1 hour. After digestion, the alleles containing a G (non SNP) produced a 450 bp fragment, while the alleles with a C (rs755622) produced a 270 bp fragment.

SNP calling. Raw BAM files from the TCGA\_GBM cohort were utilized to analyze the rs2096525 MIF SNP from whole-exome sequencing data aligned by the TCGA. The SNP rs2096525 genotype was identified via use of HaplotypeCaller, where samples with alternative counts at the reference position chr22: 23894632 were identified. After classification of the samples by genotype, the phenotype data were downloaded via TCGA, and survival analysis was performed using log-rank test via R version 4.1.0.

RNA-Seq. Flash-frozen tissue was obtained from the Rose Ella Burkhardt Brain Tumor Center at the Cleveland Clinic under IRB 2559 and corresponded to n = 34 patients with previously identified rs755622 SNP status from matched WBC pellets (n = 17 C/\*, and n = 17 G/G). Within each group, patients were selected who had undergone full Stupp protocol SOC treatment and were evenly divided by sex and prognosis (<6 months PFS or >6 months PFS). Samples were processed and sequenced by Genewiz. Briefly, RNA was extracted by Qiagen RNeasy kit, and then the library was prepared using TruSeq library preparation. Average sequencing depth was 40 Mbp per sample.

FASTQ files were aligned to the hg19 using STAR aligner with default parameters. Fragments were counted using Rsamtools with UCSC.hg19.knownGene transcript database. Raw counts were used in DESeq2 downstream for differential expression comparing the rs75662 SNP status groups. All differential expression results are provided in Supplemental Data File 1.

TCGA RNA-Seq data. Processed count data from TCGA\_GBM mRNA data set were downloaded from the Broad Firehose (https://gdac.broadinstitute.org/runs/stddata\_2016\_01\_28/data/GBM/20160128/gdac.broadinstitute.org\_GBM.mRNAseq\_Preprocess.Level\_3.2016012800.0.0.tar.gz), where the raw count file GBM.uncv2.mRNAseq\_raw\_counts.txt was utilized for downstream analysis.

Differential expression. Raw counts from TCGA\_GBM were analyzed using the R package DESeq2 version 1.29 in R version 4.0.1. After identification of the germline rs2096525 SNP status, the patient samples containing the minor allele were compared with patients homozygous for the major allele for differential expression.

Pathway enrichment. GSEA was performed based on the ranked list of differentially expressed genes by log fold-change value using the R package Gene Set Variation Analysis (GSVA). GSEA results was potted using the enrich plot R package. Hallmark gene sets used for GSEA downloaded from gsea-msigdb.

ssGSEA. The nCounter PanCancer Immune Profiling Panel gene set was used for immune infiltration deconvolution signatures. Each signature was used with ssGSEA using GSVA R package version 1.40.1. Microenvironment score, stromal score, and immune score were generated using xCell R package version 1.1.0 (59). Comparing signature scores between groups was performed using 2-way ANOVA with the P values shown for each comparison above the deconvolution heatmaps. All deconvolution and P value heatmaps were generated using pheatmap version 1.0.12.

*Immunofluorescence staining*. Serial sections (7 µm thick) from each sample (formalin-fixed paraffin-embedded tumor biopsies) were stained with 3 different sets of markers and indicated below:

Set 1 (triple immunofluorescence staining): DAPI, CD3 (ab11089, Abcam, 1:50), CD8 (85336, Cell Signaling, 1:50), CD107a (NBP2-52721, Novus Bio, 1:50); Set 2 (double immunofluorescence staining): DAPI, MIF (MAB2892, R&D systems, 1:500), LTF (HPA059976, Atlas, 1:100); and Set 3 (multiplex staining): DAPI, CD74 (ab1794, Abcam, 1:200), CD11b (ab133357, Abcam, 1:200), P2RY12 (NBP2-33870, Novus Bio, 1:200), HLA-DR (ab20181, Abcam, 1:200), CD68 (790-2931, Ventana, 1:200), CD4 (ab133616, Abcam, 1:200).



For staining, slides were baked at 60°C prior to departifinization. Slides were then deparaffinized using a Leica Autostaine XL, and antigen retrieval was performed using a sodium citrate buffer (pH 6) with slides steamed in a pressure cooker at 110°C for 10 minutes. Slides were then cooled to room temperature and transferred to water for 5 minutes prior to TBST for 15 minutes. Primary antibody was placed at the above concentrations and incubated in a humid chamber overnight at 4°C. Slides were placed on a Biocare Intellipath Staining platform for blocking (3% donkey serum), secondary antibody incubations, and Hoechst/DAPI staining (Thermo Fisher Scientific, H1399, 1 mg/mL). The following secondary antibodies were used: rabbit Cy3 (Jackson ImmunoResearch, 711-165-152, 1:250), mouse 488 (Jackson ImmunoResearch, 715-545-151, 1:250), rat Cy3 (Jackson ImmunoResearch, 712-165-153, 1:250), and rabbit Cy5 (Jackson ImmunoResearch, 711-175-152, 1:250). Slides were manually cover-slipped with an aqueous mounting medium.

Imaging. Whole-tissue sections were imaged with the multispectral capabilities of the Vectra Polaris Automated Quantitative Pathology Imaging System (Akoya Bioscience Inc.). Multispectral images were then unmixed in inForm (Akoya Biosciences Inc., version 2.5) to obtain component images for each individual marker and tissue autofluorescence. Component image tiles were stitched and saved as OME-TIFF (Open Microscopy Environment) format for analysis, storage, and archival.

Image analysis. The open-source image analysis software QuPath was used for the detection and classification of cells. For each slide, 10–20 regions of interest (ROI) were selected to represent different areas of the whole section while avoiding imaging, staining, and sectioning artifacts. StarDist (60), a deep-learning algorithm, along with a pretrained model was used within QuPath for detecting cell nuclei from the DAPI channel. For each cell, intensity measurements were used to determine its positivity for each marker in the panel. A custom script with a manual decision tree (61) was implemented in QuPath to classify cells based on their positivity. Representative images were extracted using QuPath, and single-cell data for each sample were exported as.csv files for further analysis and charting in R version 4.0.1.

NanoString GeoMx. Eight deidentified GBM samples taken from second surgeries for potentially novel MRI enhancement were evaluated using a NanoString GeoMx Digital Spatial Profiler using NanoString Human Whole Transcriptome Atlas probes. Samples were stained with Novus Sox10–Alexa Fluor 647-conjugated (catalog NBP2-59621AF647), Santa Cruz CD68–Alexa Fluor 594-conjugated (catalog sc-20060 AF594), and NanoString Syto13-conjugated (catalog S7575) antibodies to select for areas of cancer cellularity. This was confirmed with overlayed clinical IHC imaging for OLIG2, KI67, and P53. Normalization and differential expression analysis were performed using NanoString GeoMx software following the default manufacturer protocols (62–65). Clinical diagnosis of lesions was noted prior to deidentification, with 4 samples corresponding to treatment reaction and 4 samples to cancer progression.

Statistics. Demographic and clinical characteristics were evaluated between clinical cohorts. Two-way ANOVA and  $\chi^2$  tests were performed to assess differences in continuous and categorical variables, respectively. Additionally, demographic and clinical characteristics were assessed for association with the MIF SNP rs755622. For this assessment, 2-tailed t tests were performed to assess differences in continuous data. All statistics were generated in R version 4.0.1.

Overall and PFS of each clinical cohort was assessed for MIF SNP rs755622. Kaplan-Meier (KM) analysis was performed to evaluate the difference in survival and recurrence between patients with a GG genotype and with CC or CG genotypes. These analyses were also performed among only those patients who had received SOC. Log-rank tests were performed to assess differences in KM curves. Univariate and multivariable Cox proportional hazards models were generated to assess the effect of MIF SNP rs755622 on overall and PFS. The proportional hazards assumption was assessed and not found in violation. Multivariable models were adjusted for age, sex, surgery, and SOC. Hazard ratios (HR) and 95% CI are reported. All statistics were generated in R version 4.0.1.

Study approval. For Cleveland Clinic, peripheral blood samples from 451 patients with GBM were collected through the Rose Ella Burkhardt Brain Tumor and Neuro-Oncology Center under approved IRB protocol no. 2559. WBCs from each blood sample were isolated via Ficoll gradient and then snap frozen and stored at -80°C for research use. For this study, we selected all available GBM samples. For Moffitt Cancer Center, salivary DNA samples collected using Oragene kits were available for 386 recently diagnosed patients with GBM under IRB protocol no. MCC 15004. DNA was extracted and stored in aliquot pellets at -80°C for future research use. For Case Western Reserve University/University Hospitals of Cleveland, peripheral blood samples from 131 patients with GBM were collected through the Ohio Brain Tumor Study at Case Western



Reserve University, under approval from University Hospitals IRB CC296. Clinical and pathological data were gathered for each patient. Patient blood samples were collected and processed at the time of consent.

Data availability. The RNA-Seq data generated and analyzed during the current study are available in the Gene Expression Omnibus (GEO) repository under accession no. GSE232434. The data can be accessed at: https://protect-us.mimecast.com/s/i2OmCmZ6rWHprGyBlCNZghk?domain=ncbi.nlm. nih.gov. The processed data, including normalized counts and differential gene expression analyses, are also available in the supplementary materials of this manuscript (Supplemental Data File 1).

Differential expression outputs are also provided, along with summarized imaging data in the Supplemental Data File 1. All code used to generate figures can be requested from the first author TA.

#### **Author contributions**

TJA, MMG, BO, MAV, RB, TAC, MSA, and JDL conceived the experiments; TJA, MMG, BO, DB, AZ, VM, KT, MM, AR, AL, CN, GR, KAW, GC, NP, TTT, KM, and AS performed experiments and analysis; CMD, JMB, KME, CMH, JSBS, MAV, RB, TAC, MSA, and JDL supervised and participated in analysis; MAV, RB, TAC, MSA, and JDL supervised the project and provided financial support; and TJA, RB, TAC, MSA, and JDL wrote the manuscript. WW conceived experiments and performed experiments and analysis. All authors had final approval of the manuscript.

#### **Acknowledgments**

We thank Erin Mulkearns-Hubert for editorial assistance and members of the Lathia laboratory for insightful discussions. We thank also Amanda Mendelsohn for illustration assistance. This work was supported by NIH grants R01NS109742 (JDL, MAV, and RB), R35 CA232097 (TAC), K99CA248611 (DB), F31 NS101771 (TJA), and T32 CA059366 (TJA), Cleveland Clinic Research Program Committees (BO), and NIH K12CA215110 (TT) as well as the Lerner Research Institute (TAC and JDL) and Case Comprehensive Cancer Center (TAC, JMB, TAC, and JDL).

Address correspondence to: Justin D. Lathia, Lerner Research Institute, 9500 Euclid Ave, NC10, Cleveland, Ohio 44195, USA. Phone: 216.445.7475; Email: lathiaj@ccf.org. Or to: Manmeet S. Ahluwalia, 8900 N. Kendall Drive, Miami, Florida 33176, USA. Email: ManmeetA@baptisthealth.net. Or to: Timothy A. Chan, Lerner Research Institute, 9500 Euclid Ave, NB4, Cleveland, Ohio 44195, USA. Email: chant2@ccf.org.

- Verhaak RG, et al. Integrated genomic analysis identifies clinically relevant subtypes of glioblastoma characterized by abnormalities in PDGFRA, IDH1, EGFR, and NF1. Cancer Cell. 2010;17(1):98–110.
- Phillips HS, et al. Molecular subclasses of high-grade glioma predict prognosis, delineate a pattern of disease progression, and resemble stages in neurogenesis. Cancer Cell. 2006;9(3):157–173.
- Lehmann LE, et al. A MIF haplotype is associated with the outcome of patients with severe sepsis: a case control study. J Transl Med. 2009;7:100.
- 4. Xue Y, et al. The MIF -173G/C polymorphism and risk of childhood acute lymphoblastic leukemia in a Chinese population. Leuk Res. 2010;34(10):1282–1286.
- Eder L, et al. Predictors of response to intra-articular steroid injection in psoriatic arthritis. Rheumatology (Oxford). 2010;49(7):1367–1373.
- Gao L, et al. Macrophage migration inhibitory factor in acute lung injury: expression, biomarker, and associations. *Transl Res.* 2007;150(1):18–29.
- Hildebrandt MA, et al. Genetic variants in inflammation-related genes are associated with radiation-induced toxicity following treatment for non-small cell lung cancer. PLoS One. 2010;5(8):e12402.
- Llamas-Covarrubias MA, et al. Macrophage migration inhibitory factor (MIF): genetic evidence for participation in early onset and early stage rheumatoid arthritis. Cytokine. 2013;61(3):759–765.
- Falvey JD, et al. Macrophage migration inhibitory factor gene polymorphisms in inflammatory bowel disease: an association study in New Zealand Caucasians and meta-analysis. World J Gastroenterol. 2013;19(39):6656–6664.
- Wang CD, et al. Contribution of macrophage migration inhibitory factor -173G/C gene polymorphism to the risk of cancer in Chinese population. Asian Pac J Cancer Prev. 2015;16(11):4597

  –4601.
- 11. Kuai SG, et al. Functional polymorphisms in the gene encoding macrophage migration inhibitory factor (MIF) are associated with active pulmonary tuberculosis. *Infect Dis (Lond)*. 2016;48(3):222–228.
- Illescas O, et al. Macrophage migration inhibitory factor -173 G/C polymorphism: a global meta-analysis across the disease spectrum. Front Genet. 2018;9:55.
- 13. Li DY, et al. MIF -173G/C (rs755622) polymorphism modulates coronary artery disease risk: evidence from a systematic meta-analysis. *BMC Cardiovasc Disord*. 2020;20(1):300.



- Aschebrook-Kilfoy B, et al. Polymorphisms in immune function genes and non-Hodgkin lymphoma survival. J Cancer Surviv. 2012;6(1):102–114.
- 15. De la Cruz-Mosso U, et al. Macrophage migration inhibitory factor: association of -794 CATT5-8 and -173 G>C polymorphisms with TNF-α in systemic lupus erythematosus. *Hum Immunol.* 2014;75(5):433–439.
- 16. Assis DN, et al. A macrophage migration inhibitory factor polymorphism is associated with autoimmune hepatitis severity in US and Japanese patients. *Dig Dis Sci.* 2016;61(12):3506–3512.
- Castaneda-Moreno VA, et al. MIF functional polymorphisms (-794 CATT<sub>5.8</sub> and -173 G>C) are associated with MIF serum levels, severity and progression in male multiple sclerosis from western Mexican population. *J Neuroimmunol.* 2018;320:117–124.
- 18. Sharaf-Eldein M, et al. MIF-173G/C (rs755622) polymorphism as a risk factor for acute lymphoblastic leukemia development in children. *J Gene Med*. 2018;20(9):e3044.
- Banos-Hernandez CJ, et al. Macrophage migration inhibitory factor polymorphisms are a potential susceptibility marker in systemic sclerosis from southern Mexican population: association with MIF mRNA expression and cytokine profile. Clin Rheumatol. 2019;38(6):1643–1654.
- 20. Avalos-Navarro G, et al. Association of the genetic variants (-794 CATT5-8 and -173 G > C) of macrophage migration inhibitory factor (MIF) with higher soluble levels of MIF and TNFα in women with breast cancer. *J Clin Lab Anal.* 2020;34(5):e23209.
- Otvos B, et al. Cancer stem cell-secreted macrophage migration inhibitory factor stimulates myeloid derived suppressor cell function and facilitates glioblastoma immune evasion. Stem Cells. 2016;34(8):2026–2039.
- Alban TJ, et al. Global immune fingerprinting in glioblastoma patient peripheral blood reveals immune-suppression signatures associated with prognosis. JCI Insight. 2018;3(21):e122264.
- 23. Alban TJ, et al. Glioblastoma myeloid-derived suppressor cell subsets express differential macrophage migration inhibitory factor receptor profiles that can be targeted to reduce immune suppression. Front Immunol. 2020;11:1191.
- 24. Alban TJ, et al. Glioblastoma myeloid-derived suppressor cell subsets express differential macrophage migration inhibitory factor receptor profiles that can be targeted to reduce immune suppression. Front Immunol. 2020;11:1191.
- 25. Bayik D, et al. Myeloid-derived suppressor cell subsets drive glioblastoma growth in a sex-specific manner. *Cancer Discov.* 2020;10(8):1210–1225.
- Yao J, et al. Transcription factor ICBP90 regulates the MIF promoter and immune susceptibility locus. J Clin Invest. 2016;126(2):732–744.
- Melin BS, et al. Genome-wide association study of glioma subtypes identifies specific differences in genetic susceptibility to glioblastoma and non-glioblastoma tumors. Nat Genet. 2017;49(5):789–794.
- Leng L, et al. Genotyping two promoter polymorphisms in the MIF gene: a -794 CATT5-8 microsatellite repeat and a -173 G/C SNP. *Methods Mol Biol.* 2020;2080:67–84.
- 29. Zhang Y, et al. Prognostic value and risk factors of treatment-related lymphopenia in malignant glioma patients treated with chemoradiotherapy; a systematic review and meta-analysis. *Front Neurol.* 2021;12:726561.
- 30. Otvos B, et al. Preclinical modeling of surgery and steroid therapy for glioblastoma reveals changes in immunophenotype that are associated with tumor growth and outcome. *Clin Cancer Res.* 2021;27(7):2038–2049.
- Rodriguez-Franco DA, et al. [Antimicrobial mechanisms and potential clinical application of lactoferrin]. Rev Latinoam Microbiol. 2005;47(3-4):102–111.
- 32. Yamauchi K, et al. Bovine lactoferrin: benefits and mechanism of action against infections. Biochem Cell Biol. 2006;84(3):291–296.
- 33. Gonzalez-Chavez SA, et al. Lactoferrin: structure, function and applications. Int J Antimicrob Agents. 2009;33(4):301.e1–301.e8.
- 34. Zheng Y, et al. Lactoferrin suppresses the Epstein-Barr virus-induced inflammatory response by interfering with pattern recognition of TLR2 and TLR9. *Lab Invest.* 2014;94(11):1188–1199.
- Curran CS, et al. Lactoferrin activates macrophages via TLR4-dependent and -independent signaling pathways. Cell Immunol. 2006:242(1):23–30
- 36. Snyder A, et al. Genetic basis for clinical response to CTLA-4 blockade in melanoma. N Engl J Med. 2014;371(23):2189-2199.
- 37. Kang I, Bucala R. The immunobiology of MIF: function, genetics and prospects for precision medicine. *Nat Rev Rheumatol.* 2019;15(7):427–437.
- 38. Noe JT, Mitchell RA. MIF-dependent control of tumor immunity. Front Immunol. 2020;11:609948.
- Lee SH, et al. Macrophage migration inhibitory factor (MIF) inhibitor 4-IPP downregulates stemness phenotype and mesenchymal trans-differentiation after irradiation in glioblastoma multiforme. PLoS One. 2021;16(9):e0257375.
- Fukaya R, et al. MIF maintains the tumorigenic capacity of brain tumor-initiating cells by directly inhibiting p53. Cancer Res. 2016;76(9):2813–2823.
- 41. Ha W, et al. Ibudilast sensitizes glioblastoma to temozolomide by targeting macrophage migration inhibitory factor (MIF). Sci Rep. 2019;9(1):2905.
- Castro BA, et al. Macrophage migration inhibitory factor downregulation: a novel mechanism of resistance to anti-angiogenic therapy. Oncogene. 2017;36(26):3749–3759.
- 43. Meyer-Siegler KL, et al. Macrophage migration inhibitory factor (MIF) gene polymorphisms are associated with increased prostate cancer incidence. *Genes Immun.* 2007;8(8):646–652.
- Donnelly RP, et al. mTORC1-dependent metabolic reprogramming is a prerequisite for NK cell effector function. J Immunol. 2014;193(9):4477–4484.
- 45. Piette C, et al. The dexamethasone-induced inhibition of proliferation, migration, and invasion in glioma cell lines is antagonized by macrophage migration inhibitory factor (MIF) and can be enhanced by specific MIF inhibitors. *J Biol Chem.* 2009;284(47):32483–32492.
- Tuccari G, et al. Immunohistochemical detection of lactoferrin in human astrocytomas and multiforme glioblastomas. Eur J Histochem. 1999;43(4):317–322.
- 47. Arcella A, et al. In vitro and in vivo effect of human lactoferrin on glioblastoma growth. J Neurosurg. 2015;123(4):1026-1035.
- Kumari S, et al. Overcoming blood brain barrier with a dual purpose Temozolomide loaded Lactoferrin nanoparticles for combating glioma (SERP-17-12433). Sci Rep. 2017;7(1):6602.



- Liu Y, et al. Lactoferrin-induced myeloid-derived suppressor cell therapy attenuates pathologic inflammatory conditions in newborn mice. J Clin Invest. 2019;129(10):4261–4275.
- 50. Ranjan S, et al. Clinical decision making in the era of immunotherapy for high grade-glioma: report of four cases. *BMC Cancer*. 2018;18(1):239.
- Quan G, et al. Role of dynamic susceptibility contrast perfusion MRI in glioma progression evaluation. J Oncol. 2021;2021:1696387.
- 52. Gladwish A, et al. Evaluation of early imaging response criteria in glioblastoma multiforme. Radiat Oncol. 2011;6:121.
- 53. Soike MH, et al. Glioblastoma radiomics: can genomic and molecular characteristics correlate with imaging response patterns? Neuroradiology. 2018;60(10):1043–1051.
- 54. Yang I, et al. CD8+ T-cell infiltrate in newly diagnosed glioblastoma is associated with long-term survival. *J Clin Neurosci*. 2010;17(11):1381–1385.
- 55. Lohr J, et al. Effector T-cell infiltration positively impacts survival of glioblastoma patients and is impaired by tumor-derived TGF-β. Clin Cancer Res. 2011;17(13):4296–4308.
- 56. Kim YH, et al. Tumour-infiltrating T-cell subpopulations in glioblastomas. Br J Neurosurg. 2012;26(1):21–27.
- 57. Mauldin IS, et al. Proliferating CD8<sup>+</sup> T cell infiltrates are associated with improved survival in glioblastoma. *Cells*. 2021;10(12):3378.
- 58. Innocenti L, et al. The prognostic impact of gender, therapeutic strategies, molecular background, and tumor-infiltrating lymphocytes in glioblastoma: a still unsolved jigsaw. *Genes (Basel)*. 2023;14(2):501.
- 59. Aran D, et al. xCell: digitally portraying the tissue cellular heterogeneity landscape. Genome Biol. 2017;18(1):220.
- 60. Schmidt U, et al. Cell Detection with Star-convex Polygons. Paper presented at: Medical Image Computing and Computer Assisted Intervention – MICCAI 2018: 21st International Conference; September 16–20, 2018; Granada, Spain. https://link.springer.com/chapter/10.1007/978-3-030-00934-2\_30. Accessed May 26, 2023.
- 61. Viratham Pulsawatdi A, et al. A robust multiplex immunofluorescence and digital pathology workflow for the characterisation of the tumour immune microenvironment. *Mol Oncol.* 2020;14(10):2384–2402.
- 62. Beechem JM. High-plex spatially resolved RNA and protein detection using digital spatial profiling: a technology designed for immuno-oncology biomarker discovery and translational research. Methods Mol Biol. 2020;2055:563–583.
- 63. Chen Y, et al. Mapping gene expression in the spatial dimension. Small Methods. 2021;5(11):e2100722.
- 64. Hunter MV, et al. Spatially resolved transcriptomics reveals the architecture of the tumor-microenvironment interface. *Nat Commun.* 2021;12(1):6278.
- 65. Zimmerman SM, et al. Spatially resolved whole transcriptome profiling in human and mouse tissue using digital spatial profiling. Genome Res. 2022;32(10):1892–1905.