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Abstract: The role of traumatic brain injury in the development of glioma is highly controversial since first presented. This is not unexpected because traumatic brain injuries are overwhelmingly more common than glioma. However, the causes of post-traumatic glioma have been long discussed and still warrant further research. In this review, we have presented an overview of previous cohort studies and case–control studies. We have summarized the roles of microglial cells, macrophages, astrocytes, and stem cells in post-traumatic glioma formation and development, and reviewed various carcinogenic factors involved during traumatic brain injury, especially those reported in experimental studies indicating a relationship with glioma progression. Besides, traumatic brain injury and glioma share several common pathways, including inflammation and oxidative stress; however, the exact mechanism underlying this co-occurrence is yet to be discovered. In this review, we have summarized current epidemiological studies, clinical reports, pathophysiological research, as well as investigations evaluating the probable causes of co-occurrence and treatment possibilities. More efforts should be directed toward elucidating the relationship between traumatic brain injury and glioma, which could likely lead to promising pharmacological interventions towards designing therapeutic strategies.

Keywords: traumatic brain injury, glioma, promotion, mechanism

Introduction

Traumatic brain injury is a leading global cause of mortality and morbidity and the main cause of death in young people living in industrialized countries.1,2 Traumatic brain injury is mainly caused by an external mechanical force causing brain trauma. Traumatic brain injury and the ensuing neuroinflammation, in addition to causing motor and cognitive deficits, may persist long after the initial injury.3 Furthermore, long-term neuroinflammation has been related to increased risk of neurodegenerative disorders and various other deficits.4 Traumatic brain injury as a risk factor for brain tumors has been a controversial topic in medicine for over a century.5–8 However, as statistical reports on brain tumors often exclude post-traumatic glioma, relevant information on the incidence of gliomas caused by traumatic brain injuries is rare. Although previous clinical studies and case reports are often vague and difficult to evaluate since most of them were published so many years ago,5–12 some are quite striking, such as the cohort study by Munch et al in which the reduced long-term risk of malignant astrocytic tumors after structural brain injury was evaluated. However, this study was conducted using a small population. Furthermore, it is to be noted that traumatic brain injury...
injury is only one type of structural brain injury in this study. Additionally, it is challenging to confirm the incidence of post-traumatic glioma owing to the frequently considerable time gap between traumatic brain injury and glioma. Thus, there is an urgent need to systematically evaluate the role and outcome of head trauma in the incidence and progression of glioma. In this review, our primary focus is to document the interrelationship between traumatic brain injury and glioma based on a comprehensive review of the existing literature, which is discussed in detail. First, we present an overview of previous cohort studies and various case reports regarding the relationship between traumatic brain injury and glioma. Next, we discuss the roles of microglial cells, macrophages, astrocytes, and stem cells in post-traumatic glioma formation and development. Moreover, we also briefly discuss the various carcinogenic factors during traumatic brain injury that could explain the interplay between these two parameters. We have also summarized the common inflammatory and oxidative stress-related signaling pathways related to traumatic brain injury and glioma. Lastly, we have elaborated on the strategy that could be considered in a clinical setting, and have concluded this review with directions for future research.

An Overview of Previous Epidemiological Studies

All previously published cohort studies and case-control studies are not directly comparable owing to differences in exposures and outcomes (Table 1). A population-based study by Inskip et al. reveals an increased overall incidence of intracranial tumors of head trauma patients, whereas no significant association was found in the case of malignant astrocytic tumors. In a cohort study by Nygren et al. no significant association between traumatic brain injuries and brain tumors was identified; moreover,
Specific risks for malignant astrocytic tumors were not reported. However, more recently, a cohort study by Chen et al. indicated an increased risk for not otherwise specified malignant brain tumors within 3 years after a traumatic brain injury. Besides, interestingly, a research group demonstrated a decreased risk 5 or more years after structural brain injury; however, they did not find convincing evidence for an association between structural brain injury and malignant astrocytic tumors within the first 5 years of follow-up. The authors speculated that the inflammatory response after traumatic injury could cause elevated immunological alertness for astrocytes undergoing neoplastic transformation, as well as a clearance of premalignant astrocytes or neural stem cells, which may otherwise have developed into glioma. Although their study demonstrated that structural brain injury may generally reduce the long-term risk of malignant astrocytic tumors, their data also supported that structural brain injury specifically caused by trauma (different from other types of exposures such as cerebral ischemic infarction and intracerebral hemorrhage) could increase the long-term risk of malignant astrocytic tumors. Thus, the relationship between traumatic brain injury and glioma is still not conclusive and warrants further studies.

### Table 1 Overview of Published Epidemiological Studies Exploring the Causal Relationship Between Traumatic Brain Injury and Glioma

<table>
<thead>
<tr>
<th>Series</th>
<th>Study Design</th>
<th>Region</th>
<th>Age</th>
<th>Cohort Study: Participants; Glioma Cases; Mean Follow-Up</th>
<th>Case-Control Study: Cases/Controls</th>
<th>Data Collection</th>
<th>Risk Estimates</th>
</tr>
</thead>
<tbody>
<tr>
<td>Inskip 1998</td>
<td>Cohort</td>
<td>Denmark</td>
<td>22.5 (Mean)</td>
<td>224,421 patients; 79 glioma cases; Mean follow-up 8 years</td>
<td>Follow up</td>
<td>SIR, 1.0 (0.80–1.20)</td>
<td></td>
</tr>
<tr>
<td>Nygren 2001</td>
<td>Cohort</td>
<td>Sweeden</td>
<td>32.2 (Mean)</td>
<td>311,006 patients; 161 glioma cases; Mean follow-up 10.4 years</td>
<td>Follow up</td>
<td>SIR, 1.0 (0.90–1.20)</td>
<td></td>
</tr>
<tr>
<td>Chen 2012</td>
<td>Cohort</td>
<td>Taiwan</td>
<td>43.5 (Mean)</td>
<td>5007 patients; 9 glioma cases; Mean follow-up 3 years</td>
<td>Follow up</td>
<td>SIR, 2.08 (1.27–3.22)</td>
<td></td>
</tr>
<tr>
<td>Munch 2015</td>
<td>Cohort</td>
<td>Denmark</td>
<td>51 (Mean)</td>
<td>48,194 patients; 14 glioma cases; Median follow-up 28 years</td>
<td>Follow up</td>
<td>RR, 1.99 (1.00–3.50)</td>
<td></td>
</tr>
<tr>
<td>Hochberg 1984</td>
<td>Case-control</td>
<td>USA</td>
<td>15–81</td>
<td>125/113</td>
<td>Questionnaire</td>
<td>&lt;15yrs: RR, 1.1 (0.40–3.20) for mild trauma; RR, 1.2 (0.30–5.60) for severe trauma; ≥15yrs: RR, 1.4 (0.40–4.70) for mild trauma; RR, 10.6 (2.10–53.30) for severe trauma</td>
<td></td>
</tr>
<tr>
<td>Zampieri 1994</td>
<td>Case-control</td>
<td>Italy</td>
<td>18–70</td>
<td>195/195</td>
<td>Questionnaire</td>
<td>OR, 0.7 (0.30–1.40) for all glioma types; OR, 1.2 (0.30–5.00) for low-grade glioma; OR, 0.5 (0.20–1.30) for high-grade glioma</td>
<td></td>
</tr>
<tr>
<td>Hu 1998</td>
<td>Case-control</td>
<td>Northeast China</td>
<td>Not found</td>
<td>218/436</td>
<td>In-person interview or questionnaire</td>
<td>OR, 4.09 (2.51–10.31)</td>
<td></td>
</tr>
<tr>
<td>Preston-Martin 1998</td>
<td>Case-control</td>
<td>Six countries</td>
<td>20–79</td>
<td>1178/1987</td>
<td>Questionnaire</td>
<td>Male: 1.18 (0.94–1.48) for any trauma; 1.13 (0.87–1.48) for serious trauma; Female: 1.03 (0.42–2.55) for any trauma; 1.07 (0.74–1.56) for serious trauma</td>
<td></td>
</tr>
</tbody>
</table>

**Abbreviations:** SIR, standardized incidence ratio; RR, relative risk; OR, odds ratio.
It is often a challenge to compare the results of previously published epidemiological studies, as it involves individuals of different ages who live in different environments. Importantly, most studies have also not been standardized regarding the type or the severity of brain damage. The low incidence of brain tumors also hinders the design of relevant research. There is currently an urgent need for more comprehensive and larger-scale epidemiological investigations, including cohort studies and case-control studies, to evaluate post-traumatic glioma. To date, very few case-control studies have specifically reported the risk for malignant astrocytoma/glioma after traumatic brain injury, and conclusively, the currently available findings are equivocal with null or positive associations. For the first time, Hochberg et al did a case-control study of 160 persons with glioblastoma, and the results suggested that severe head trauma in adults is a significant risk factor for glioblastoma. After that, Zampieri et al did another case-control study to find potential risk factors for cerebral glioma in adults, however, their study yielded no meaningful association between head trauma and glioma. Besides, the case-control study done by Preston-Martin et al investigated the role of head trauma from injury in adult brain tumor risk. Although not significant association between head trauma and glioma has been found, their findings suggest that an association between head trauma and brain tumor risk cannot be ruled out and should therefore be further studied, and future studies of head trauma and brain tumor risk should consider potential initiators of carcinogenesis, such as nitrite from cured meats, as modifiers of the trauma effect on risk of brain tumor. Furthermore, Hu et al also exerted case-control study of risk factors for glioma in adults, interestingly positive associations between brain trauma and glioma has been found. Unfortunately, all case-control studies were conducted before the year 1998, and no newly published research worthy of reference could be found suitable for discussion in this review. The advantages of cohort studies have been highlighted in various studies; therefore, most researchers give more weightage to cohort studies than case-control studies when systematically evaluating evidence. Thus, additional more cohort investigations with correct and standardized study designs are much needed to gain a better understanding of post-traumatic glioma.

An Overview of Previous Case Reports

The results of the published epidemiological studies could not be compared uncritically, as the types of brain injuries differed, and patients belonged to different ethnic groups and different ages. Difficulties in conducting epidemiological studies can be attributed to the low incidence of brain tumors. More efforts should be directed toward investigating the causal relationship between traumatic brain injury and glioma, which is supported by several published case reports. Although these reports from epidemiological observations have not conclusively confirmed the relationship between traumatic brain injury and glioma, some reports discuss the follow-up details of patients in great detail and are indicative of the possibility of such a relationship. Anselmi et al reported two cases of brain glioma that developed in the scar of an old brain trauma, these two cases fulfill the established criteria for a traumatic origin of brain tumors and add further support to the relationship between cranial trauma and glioma. Di Trapani et al reported that several years after sustaining a commotio left parietal trauma, one patient developed a mixed glioma in the left temporo-parietal-occipital region in continuity with the scar resulting from the trauma. Magnavita et al reported the case of a patient who suffered a severe head injury to the right temporoparietal lobe, and the patient developed a glioblastoma multiforme at the precise site of the meningocerebral scar 4 years later. Moorthy et al reported a case of a 56-year-old man who had history of head injury 5 years prior with CT evidence of bilateral basifrontal contusions. Imaging showed a large left frontal intra-axial mass lesion and the histopathology was reported as glioblastoma multiforme. The authors formulated additional radiologic criteria for tumors that may present following trauma. Mrowka et al reported that a glioblastoma multiforme developed 30 years after a penetrating cranioencephral injury in the left parietal region caused by fragments of an artillery projectile. Sabel et al reported that a patient developed a left-sided frontal glioblastoma multiforme at the precise site of the meningocerebral scar and posttraumatic defect 37 years later. Witzmann et al reported a case of a 28-year-old male who suffered a frontal penetrating gunshot injury with subsequent bifrontal brain abscess and subdural empyema, and five years later developed a large bifrontal glioblastoma multiforme at the precise site of the meningo-cerebral scar and posttraumatic defect. Zhou...
et al.\textsuperscript{22} also reported one case of glioblastoma multiforme that developed in the scar of an old brain trauma 10 years ago. Han et al.\textsuperscript{23} presented the first case of pregnancy-related post-traumatic malignant glioma in a 29-year-old female, and suggested that pregnancy may promote the manifestation of the clinical symptoms. Tyagi et al.\textsuperscript{24} used radiographic evidence from two patients to assess the possibility of a link between TBI and glioblastoma multiforme. Salvati et al.\textsuperscript{25} presented 4 cases of post-traumatic glioma with radiological evidence of absence of tumor at the time of the injury. Henry et al.\textsuperscript{26} reported a case of post-traumatic malignant glioma with radiological evidence of only a contusion prior to the development of the glioma. Simińska et al.\textsuperscript{27} reported one case of post-traumatic glioma 2 years after head injury. Overall, some data from these studies might support the conclusion that the association is almost weak, while others not; but a causal relationship between traumatic brain injury and glioma is highly possible. This is because traumatic brain injury initiates inflammation, oxidative stress, repair, oncogene activation, and other pathophysiological changes, which are bound to lead to malignancy in at least some patients.\textsuperscript{28,29}

Besides, to better identify reported cases addressing the relationship between traumatic brain injury and the incidence and development of glioma, an important aspect is to be able to recognize and differentiate between a tumor, traumatic brain injury-induced glioma, and post-traumatic glioma. We believe that only specific cases that fulfill certain conditions or criteria, could add to revealing the etiological association between head trauma and glioma. Thus, more efforts should be directed in establishing if these criteria, as well as possibly other cases with accurate retrospective data of traumatic brain injury and high risk of developing glioma, could add to the clarification of the etiological association between traumatic brain injury and glioma.

**Role of Microglial Cells**

Neuroinflammation accompanying the activation of microglial cells and other effector cells has been suggested as an important mechanism of TBI.\textsuperscript{33} Active microglial cells can transform to the M1 phenotype, to secrete proinflammatory or cytotoxic mediators that mediate post-TBI cell death and neuronal dysfunction, or to the M2 phenotype, to participate in phagocytosis and secrete anti-inflammatory cytokines and neurotrophic factors that are important for neural protection and repair.\textsuperscript{34} Indeed, they can become polarized ranging from the classic M1 phenotype to an alternative M2 phenotype after TBI.\textsuperscript{35} The M1 response is presumed to be pro-inflammatory,\textsuperscript{36} whereas the M2 phenotype owns anti-inflammatory effects.\textsuperscript{37} Multiple molecular pathways, such as STAT, nuclear factor-kB (NF-kB), and interferon regulatory factor (IRF), are involved in the regulation of M1/M2 phenotypic transitions.\textsuperscript{38–40} Preclinical evidence indicated that mixed phenotypes are present in the pathological processes of TBI, which offer opportunities for therapeutic interventions.\textsuperscript{41}

Several mechanisms have been shown to be associated with the formation of post-traumatic glioma, specifically, inflammatory processes and oxidative stress, both of which are mainly involved in the removal of damaged components from the brain and are known to play irreplaceable roles in this process.\textsuperscript{24} In the brain, these mechanisms are mainly mediated by the microglia or other cells of the immune system.\textsuperscript{24,42} Microglia in the brain play a role in phagocytosis and antigen presentation, leading to the release of chemokines or cytokines.\textsuperscript{43} Interestingly, recent in vivo studies have shown that microglia could have different effects on the development of brain glioma, and also result in immunosuppressive conditions that promote glioma development.\textsuperscript{44,45} Although the growth-promoting effect of glioma by microglia after traumatic brain injury is controversial, its significant role in promoting an environment that can facilitate glioma development has been identified.\textsuperscript{43} Microglia can produce metalloproteinases in the tissues adjacent to glioma, which can facilitate tumor invasion.\textsuperscript{44} Besides, PGE2 can also contribute to the creation of an environment that facilitates glioma development.\textsuperscript{46} PGE2 is released by the microglia accompanying the developing glioma and can suppress T lymphocytes. The net effect is a decreased expression of major histocompatibility complex (MHC) class II molecules on antigen-presenting cells.\textsuperscript{46} Brain-repair processes
mainly involve the microglia in normal conditions; however, during traumatic brain injury, various other cells from the immune system can also enter the brain parenchyma along with blood. These effects cannot be ignored.

Role of Macrophages

Oxidative stress caused by ROS in the acute phase of TBI and cerebral infarction is thought to be detrimental, and macrophages have been recognized as the main cells that produce ROS. During traumatic brain injury, macrophages migrate to the site of the damaged blood-brain barrier and secrete interleukin 6 (IL-6). In normal conditions, the expression of IL-6 is very low, whereas, during traumatic brain injury, its production increases considerably. Brain injury elevates IL-6 production in both serum and CSF to high concentration. Notably, multiple TBI patients’ samples have also showed that the combination of elevated IL-6 concentrations is correlated with better outcomes in patients with TBI, suggesting IL-6 as a new therapeutic strategy as well as for prediction of disease outcome of patients with TBI. Importantly, high levels of IL-6 in the brain generally result in an adverse impact on microcirculation and lead to the destruction of the blood-brain barrier in an obviously wider area compared to the initial area of trauma. Thus, in traumatic brain injuries, it is crucial that the blood-brain barrier is not initially compromised, as IL-6 can subsequently promote the entry of macrophages to the site of injury and aggravate brain edema. Besides, Xu et al reported that IL-6 also impacts cell-cycle regulation and activates signal transducer and activator of transcription-3 (STAT3), which is important for cell proliferation, differentiation, and apoptosis. Previous studies show that STAT3 inhibition suppresses the growth of glioma cells and promotes apoptosis. These findings have also been confirmed in other in vivo studies. Besides, STAT3 activation can inhibit T lymphocytes and MHC II molecules on microglial and other antigen-presenting cells. Thus, STAT3 has an immunosuppressive effect and is likely a carcinogenic factor for glioma. Importantly, the increased concentrations of IL-6 and its receptors in the cerebrospinal fluid of patients who underwent traumatic brain injury are indicative of the involvement of IL-6 in glioma progression.

Role of Neuronal Stem Cells

The neuronal stem cells in the brain are mainly generated from the subgranular zone of the hippocampal dentate gyrus and the subependymal zones of the lateral ventricles. Traumatic brain injury leads to the migration of neuronal stem cells to the damaged sites to promote regeneration, thereby differentiating into astrocytes, neurons, and oligodendrocytes. Additionally, neuronal stem cells could release cytokines and neurotrophic factors such as glial cell-derived neurotrophic factor (GDNF) and brain-derived neurotrophic factor (BDNF). Thus neuronal stem cells may be an effective treatment for neurological recovery after TBI. Interestingly, neuronal stem cells show a high expression of oncogenic genes and high sensitivity to chemical mutagenic factors. This is important because stem cells are involved in the production of ROS and various pro-inflammatory factors. Neuronal stem cells can be highly sensitive to mutagenic agents and could, thus, be easily mutated as a result of the action of certain agents. These characteristics may promote the formation of rapidly proliferating tumor cells and increase the progression of glioma in the brain. Migration of stem cells has already been identified in cases of traumatic brain injury, ischemia, and demyelination. However, there is a much higher risk of increased neoplastic transformation only during traumatic brain injury. Therefore, it is reasonable to accept the causal relationship between traumatic brain injury that induces brain stem cell activity and subsequent development of glioma. Stem cells have been generally recognized as potentially oncogenic in glioma and several studies have demonstrated their important role in the formation of gliomas. The role of stem cells should be emphasized in the analysis of relevant mechanisms leading to glioma development induced by traumatic brain injury.

Role of Astrocytes

Multiple studies have suggested that astrocytes play a key role in the pathogenesis of TBI. Increased reactive astrocytes and astrocyte-derived factors are generally observed in both experimental animal models and TBI patients. Astrocytes have beneficial and detrimental effects on TBI, including acceleration and suppression of neuroinflammation, promotion and restriction of neurogenesis and synaptogenesis, and disruption and repair of the BBB through various bioactive factors. Additionally, astrocytic aquaporin-4 is also involved in the formation of cytotoxic edema. Thus, astrocytes are attractive targets for novel therapeutic drugs for TBI. Based on case reports studying glioblastomas, neoplastic transformation of damaged astrocytes has been proposed as a possible mechanism occurring at the site of traumatic brain injuries.
accepted that astrocytes are essential components of the blood-brain barrier, and damage to the blood-brain barrier often occurs after the action of pro-inflammatory prostaglandins and leukotrienes, which triggers the effect of the relaxing of tight junctions. Pro-inflammatory factors lead to a relaxation of the capillary epithelium and the glial cells are exposed to potentially mutagenic agents. Traumatic brain injury accompanied by damage to the blood-brain barrier always causes a recovery reaction, which explains the recurrence of glioma in some cases.

To conclude, the summary of various carcinogenic factors that play a role during traumatic brain injury is presented in Figure 1. Traumatic brain injury can lead to the influx of macrophages to the site of brain injury, where they are activated and produce IL-6. Traumatic brain injury also induces enhanced IL-6 secretion by astrocytes and microglial cells. Increased IL-6 levels caused by traumatic brain injury can thus activate STAT3, thereby increasing cell proliferation at the site of injury and resulting in the inhibition of apoptosis. STAT3 can suppress T lymphocytes and decrease the activity of MHC class II molecules on cells of the immune system. Furthermore, the increased levels of IL-6 also impact the blood-brain barrier. Additionally, microglial cells secrete metalloproteinases in the tissues adjacent to the tumor, facilitating its migration and, consequently, facilitating its development. PGE2, which is synthesized by microglial cells during the development of the glioma, suppresses the T lymphocytes and decreases the expression of MHC II molecules. Besides, the generation of reactive oxygen species (ROS) might lead to certain mutations in stem cells that migrate to the injury site. At the site of injury, the risk of mutations and cell proliferation increases, and along with the inhibition of apoptosis, these factors may jointly contribute to carcinogenesis.

**Common Inflammatory and Oxidative Stress Signaling Pathways in Traumatic Brain Injury and Glioma**

Inflammation at the site of traumatic brain injury and glioma has been well documented in the literature. Besides, ROS, the major contributor of oxidative stress, are metabolic byproducts originating from different sources in hypoxic conditions and exhibit condition-dependent functions. The activation of inflammation and oxidative stress are reported in both traumatic brain injury and glioma, and both conditions appear to share a common network of signaling for downstream functions (Figure 2). Interestingly, the activation of inflammation can also contribute to oncogenesis via the generation of ROS and the activation of oxidative stress, and conversely, oxidative stress also promotes inflammation. Specifically, in this situation, astrocytes, microglia, stem cells, and even neurons can be stimulated to increase ROS and RNS (NO, ONOO−), which participate in regulating inflammation and oxidative stress in traumatic brain injury and glioma.

After traumatic brain injury, there is sequential migration of the resident microglia and myeloid inflammatory cells to the site of injury. These inflammatory cells contribute to oncogenesis via promoting ROS generation, which has mutagenic properties, or via the secretion of cytokines and growth factors, in addition to maintaining an inflammatory response. During oxidative stress or inflammation in the brain, there is an increase in ROS that can generation in the mitochondria. Several cancer-specific external stimuli, including TNF-α, lead to an increase in the mitochondrial membrane potential and interfere with the components of the electron transport chain (ETC), thereby promoting ROS generation. Besides, the NADPH oxidase (NOXs) protein family is one of the main producers of ROS in various cancers and traumatic brain injuries. Moreover, specific markers, such as TGF-β, MAPK, AKT, and ERK, among others, can lead to conformational changes in the NOX complex and increase ROS generation. Another indispensable pathway that has an impact on glioma and traumatic brain injury is hypoxia-inducing factor 1 (HIF-1), which can be upregulated owing to the inhibition of degradation via the inactivation of PHD. HIF-1 increases the expression of glucose transporter 3 (GLUT3), erythropoietin (EPO), VEGF, and BNIP3. Several other signaling pathways are involved in the activation of inflammation and oxidative stress. Nuclear factor-κB (NF-κB) can increase ROS generation via a positive feedback loop of TNF regulation. Additionally, ROS can be regulated by the Ras-Raf-MEK pathway through the transcriptional regulation of GATA-6. It has been reported that transcriptional enhancement of HSF1 by Ras upregulates SESN1 and SESN3 genes to promote ROS production. Besides, TGFβ also increases the production of ROS by activating the GSK3β and mTOR signaling pathways in the mitochondria and inhibiting antioxidant enzymes, including SOD and glutathione peroxidase (GPx) (Figure 2).
Figure 1 Schematic representation of various carcinogenic factors during traumatic brain injury. Traumatic brain injury could lead to the migration of macrophages to the site of injury, followed by increased IL-6 production. Traumatic brain injury also induces enhanced IL-6 secretion by astrocytes and microglial cells. The increased IL-6 could thus activate STAT3, which increases cell proliferation at the site of injury, as well as inhibition of apoptosis. STAT3 suppresses T lymphocytes and inhibits major histocompatibility complex (MHC) class II molecules on cells of the immune system. The increased IL-6 also damages the blood-brain barrier (BBB). In addition, microglia secretes metalloproteinases in the tissues adjacent to the tumor, facilitating its migration and development. PGE2 is also synthesized by microglia and suppress T lymphocytes, and also decrease the expression of MHC II molecules on antigen-presenting cells. Besides, reactive oxygen species (ROS) might lead to certain mutations in stem cells that migrate to the site of injury. At the site of injury, the risk of these mutations, and cell proliferation increase, as well as the apoptosis inhibition, may jointly contribute to carcinogenesis.
Clinical Approach and Future Research

Following a traumatic brain injury, there is an increase in free radicals and the expression of several pro-inflammatory genes by various transcription factors such as NF-κB. This knowledge could be used in anticancer drug discovery. ROS levels increased by "oxidation therapy" can trigger cell death via necrosis or apoptosis. Flavonoids, such as quercetin, catechins, and proanthocyanins, protect glial cells from inflammation and oxidative stress. These compounds exert protective effects in the brains of patients with cancer and help prevent traumatic brain injury. An anticancer agent, gallic acid, is not only toxic to glioma cells but also exerts beneficial effects in the recovery from traumatic brain injuries. Cardamonin (a chalcone) is effective as an anti-inflammatory and anti-carcinogenic agent in glioma. Hyperbaric oxygen (HBO) therapy is a recently developed method that has been extensively used as an adjuvant in the treatment of various diseases predominantly related to hypoxic conditions. As traumatic brain injury and glioma are related to hypoxia, HBO therapy may be expected to be efficacious in the management of these diseases. However, there could be significant differences in outcomes among patients, depending on the size of the lesion, tumor type, and malignancy. Besides, several drugs, including glycyrrhizin,
salidroside,116–118 and astragaloside,119,120 may be used in both glioma and traumatic brain injury treatment due to the counteracting effect of common signaling pathways (Figure 3).

Currently, comprehensive research establishing the relationship between the mechanisms of traumatic brain injury and tumorigenesis is necessary. However, there could be some obstacles. First, the considerable time interval between brain injury and the onset of glioma poses a challenge to perform in vivo studies. Secondly, designing in vitro studies using primary cultures can also be difficult owing to a large number of different types of cells that constitute the brain tissue. The use of immortalized glial cell lines is also excluded owing to their physiological dissimilarity with normal brain cells. Therefore, more efforts should be directed toward establishing suitable in vivo and in vitro models to explore the causal relationship between traumatic brain injury and glioma.

**Conclusion**

The possible association between traumatic brain injury and glioma should be further examined by designing additional experimental and clinical research. Much more additional factors may be involved in the formation of the post-traumatic glioma. These factors might have been unintentionally omitted during the selection of study groups in various
previous studies, leading to the result of the lack of connection between injury and glioma, which is why further explorations on the etiology of post-traumatic glioma are urgently needed. Besides, it may be more difficult to effectively treat patients who suffer from both glioma and traumatic brain injury compared to those with traumatic brain injury without glioma. The survival rate of patients with glioma is bound to increase with the development of anticancer drugs, including those suggested in this review. Treating traumatic brain injury in patients with glioma can be still challenging and requires specific treatment modalities. Thus, the development of effective strategies in the management of traumatic brain injury in patients with glioma is essential.

Copyright/Ethics Statement

The authors warrant that the article and all figures included in this work are the authors’ original work and has not been published before.

Disclosure

The authors declare no competing financial interests and no conflicts of interest for this work.

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