Dysregulation of Non-Heme Iron Metabolism in Glial Brain Tumors

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\begin{abstract}

\textbf{Objective:} To study iron exchange irregularities in experimental animals and patients with glial brain tumors to ascertain the role of the ‘iron component’ in glial brain tumor pathogenesis. \textbf{Subject and Methods:} A suspension of A.101.08 tumor cells was implanted in the cortex of the left-brain hemisphere of rats to model experimentally induced glial brain tumor. At 7 or 14 days after implantation, blood and tissue samples from the tumor, peritumoral tissue, and brain regions were taken for analysis. Blood and plasma samples were obtained from 23 patients as well as biopsy samples taken during tumor removal surgery. Electron resonance spectroscopy was used to determine the concentrations of transferrin iron, transferrin in whole blood and in blood cells, and chelatable and stored iron in the tissues of experimental animals and patients. \textbf{Results:} Hypoferremia was found in rats with both small and large glial brain tumors, whereas hyperferremia was found to be a characteristic of malignant glial brain tumors in humans. We identified statistically significant increases in stored and chelatable iron concentrations in the tumor and peritumoral brain tissue compared to the blood and the adjacent brain tissue (probably normal) in both human malignant glial brain tumors and in rat experimental glial brain tumors. \textbf{Conclusions:} These findings suggest that iron misregulation plays a part in glial brain tumor pathogenesis and this may provide a basis for understanding the association between glial brain tumors and epilepsy.

\end{abstract}

\section*{Introduction}

Epidemiological studies point to the association of iron overload with liver carcinoma, colorectal and ovarian cancers [1–3]. Individuals who are homo- or heterozygous for hereditary hemochromatosis also have an increased risk of developing neoplastic diseases [4]. Iron overload has been shown to be a risk factor for cancer development in animal models whereas iron depletion prevents the growth of cancer cells in vitro [5]. Iron ‘withholding’, resulting in hypoferremia, is considered to be a defense against neoplasia [6], and hypoferremia is characteristic of carcinoma, melanoma, sarcoma and cancers of the hematopoietic tissues in humans [7]. Increased concentrations of low-molecular weight iron (LMWI) complexes are considered to contribute significantly to...
tissue damage through such mechanisms as free radical reaction catalysis, DNA injury, a shift to cell proliferation rather than apoptosis, and inhibition of mitochondrial electron transport [8–10]. Brain tissues are especially susceptible to oxidative damage due to the high content of unsaturated fatty acids and relatively low activity of the antioxidant system compared to other tissues [11].

Tumor cells, including brain tumor cells and especially glioblastomas, are characterized by increased transferrin receptor expression compared to normal brain tissues, both in vitro and in vivo [9]. While ferritin synthesis is activated in some cancer cells [12], much less is known about the shifts in non-heme iron exchange in brain tumor growth. Hence we decided to investigate non-heme iron exchange in experimental rat glial brain tumors and in biopsied human glioblast tumors using an electron spin resonance (ESR) technique to determine if the non-heme iron composition of the blood and brain regions were linked to brain tumor pathogenesis.

**Subjects and Methods**

**Experimental Rat Model of Gliial Brain Tumor Formation**

The animal research was approved by the Ethics Committee, Romodanov Institute of Neurosurgery, Academy of Medical Sciences, Ukraine (AMSU). A suspension of A.101.08 tumor cells (atyypical astrocytoma, close to glioblastoma) was implanted into the cortex of the left-brain hemisphere of male rats aged 7–8 days (n = 14). At 7 or 14 days following implantation, samples of whole heparinized blood and liver tissue as well as samples of the brain tissue from the tumor (perifocal zone, cortex zones, the right hemisphere cortex, and hypothalamus) were removed for analysis. Samples of whole heparinized blood, liver, cortex and hypothalamus were also taken from the control animals (males, aged 18–20 days, n = 5). Animals with glial brain tumors were divided into two groups according to the size of the tumor: group 1 rats with a tumor diameter less than 3 mm (small tumors, n = 8) and group 2 rats with a tumor diameter greater than 8 mm (large tumors, n = 6).

**Patients**

The study protocol was also approved by the Ethics Committee, Romodanov Institute of Neurosurgery, A.M.S.U. A total of 40 subjects were studied: 23 patients with glial brain tumors (17 men and 6 women, mean age 39.5 ± 3.5 years) and 17 control subjects (8 men and 9 women, mean age 42.6 ± 2.1) without tumors. Of the 23 patients, 8 had grade III–IV anaplastic astrocytoma, 3 grade III oligodendroastrocytoma, 2 grade III–IV ependymoastrocytoma, 1 protoplasmic astrocytoma, 1 fibrillary astrocytoma, and 5 glioblastoma. In the morning on the day before surgery, venous blood samples were collected in heparinized tubes from overnight fasted patients and control subjects. Plasma was recovered by centrifugation of the heparinized blood at 1,300 g for 10 min. The packed cell volume, or hematocrit (Ht), was determined by centrifuging the blood specimen in capillary tubes and measuring the relative height of the red blood cell column. Biopsy samples were taken from three brain regions after radical removal of the tumor: the tumor zone, the perifocal zone adjacent to the tumor (peritumoral zone I), and the perifocal zone distant from the tumor (peritumoral zone II).

**Chemicals and Solutions**

Human apo-transferrin (product No. T2252), HEPES sodium salt (H7006), deferoxamine (D9533), iron (III) citrate (F61290), tris(hydroxymethyl)aminomethane (T1503), ammonium iron (III) sulfate (F1018), thioglycolic acid sodium salt (T0632), and 1,10-phenanthroline monohydrate (P9375) were purchased from Sigma-Aldrich (St. Louis, Mo., USA).

All chemical operations were carried out using double distilled deionized water. To remove all traces of metals, glassware was washed with 1% NaOH and 2% EDTA aqueous solution. To prepare transferrin iron calibration samples, aliquots of 25 μl iron (III) citrate in the range of 0.7–7 mM concentration were incubated with 150 μl of 0.24 mM apo-transferrin in 1 M HEPES buffer, pH = 7.4, in the presence of excess carbonate (final concentration 20 mM) for 24 h at 4°C. These were then frozen in liquid nitrogen and stored until used.

To prepare iron (III)-Df calibration samples, 25 μl of ammonium iron (III) sulfate solution with an iron concentration in the range of 1.5–15 mM was incubated with 100 μl of 1.5 mM deferoxamine B solution in 0.5 M Tris-HCl buffer, pH = 7.0, for 10 min, frozen in liquid nitrogen and stored until used.

The iron concentration in the stock solutions was determined using a colorimetric method with 1,10-phenanthroline [13].

**Treatment of Tissue Samples**

Blood, plasma, and tissue samples for ESR analysis (8- to 100-mg microsamples) were packed into cryoresistant, calibrated polymer ampoules and quickly frozen in liquid nitrogen (77 K), where they were stored until measured. Storage of the samples in liquid nitrogen did not have any discernable effect on the ESR spectra.

To determine the transferrin protein concentration, 75–100 μl blood samples were treated with 20 μl of 2.5 mM Fe (III) citrate and 20 mM sodium bicarbonate in 0.1 M HEPES, pH 7.4, at 4°C for 24 h, frozen and stored until ESR measurements were made. To determine the amount of iron available for chelation, up to 100 mg of blood or tissue samples were mixed with 25 μl of 15 mM Df solution in 0.5 M Tris-HCl buffer, pH 7.0, incubated for 30 min at 37°C, frozen and stored until ESR measurements were made.

**ESR Spectra Registration and Calculations**

The ESR spectra were obtained using an EPA-10 mini spectrometer (St. Petersburg Instruments, Russia), at super high frequency of 9.30 GHz, on pressed cylindrical samples. The ESR spectra were recorded under nonsaturating conditions.

The transferrin iron concentration was determined using characteristic iron (III)-transferrin complex spectra at an effective g-factor of 4.3 (fig. 1, curve a) [14]. The chelatable iron concentrations in tissue and blood samples were determined based on the incorporation of LMWI into the iron-deferoxamine B (Df) complexes with a characteristic ESR spectrum (fig. 1, curve b) upon treatment with Df solution [15, 16]. The stored iron concentration in tissues was evaluated against the background of the wide magnetic resonance spectra in the g-factor range of 2.0–4.5 stipulated by ferritin and/or hemosiderin iron [17–19] (fig. 2). The relative methemoglobin (MtHb) iron concentration was estimated using an amplitude...
of the sharp ESR absorption derivative at \( g = 6.0 \), corresponding to the high-spin (d^5) Fe(III) configuration of the ferric iron in oxidized hemoglobin [20].

A CuEDTA sample of the same shape and size as above, prepared in a 1:1 water-glycerin matrix, was used as a reference sample. The g value measurements were performed using the sextet of the Mn^{2+} in MgO matrix as a secondary standard. To improve the accuracy of the double integration, the ESR lines were simulated as a sum of Lorentzian profile derivatives after subtraction of the background signal, as shown in figure 1, curves a’ and b’.

Calibration curves of the double integrated intensity versus iron concentration were obtained for each of the iron species. The tissue ferritin ESR spectra were calibrated against Mössbauer data [17].

The concentration of transferrin [Tf] in the blood or plasma was calculated as [Tf] = [Tf-Fe]_{sat}/2, where [Tf-Fe]_{sat} was the transferrin iron content in a sample saturated with iron. Transferrin saturation with iron was calculated as \%Tf = 100% × [Tf-Fe]/[Tf]. The concentration of transferrin iron associated with blood cells [Tf-Fe]_{bc}...
was calculated using the transferrin iron concentration in the blood \([\text{Tf-Fe}]_{\text{bl}}\) and plasma \([\text{Tf-Fe}]_{\text{pl}}\), and accounting for the \(\text{Ht}\) as follows:

\[
[\text{Tf-Fe}]_{\text{bc}} = \frac{[\text{Tf-Fe}]_{\text{bl}} - [\text{Tf-Fe}]_{\text{pl}} \times (1 - \text{Ht})}{\text{Ht}}
\]

### Statistical Analysis

Data are represented as mean ± SEM, where SEM is the standard error of the mean. To determine the difference between experimental and control data sets, the quantitative statistical measure, \(P_1\) (distance between data sets), was used: \(P_1 = \Phi_{nm}(T_1(x, y))\), where \(\Phi_{nm}\) is the Snedecor-Fischer function:

\[
T_1(\xi, \eta) = \frac{n + m - I - 1}{I} + \frac{nm}{(n + m)(n + m - 2)} D^2_s(\xi, \eta)
\]

where \(D^2_s(\xi, \eta)\) is a Mahalanobis distance, and \(I\) is the number of parameters (factors). Parameters \(n\) and \(m\) were determined by the sample size for experimental and control data. The advantages of this estimation are the following: the \(P_1\) value is a normalized one and varies in the range of 0–1; the closer \(P_1\) is to the unit, the less probable is the so-called ‘zero’ hypothesis on the absence of the effect. This estimation takes into account the interrelationships between factors and their coupled changes. In the partial case of \(I = 1\), \(P_1 = S_{nm}(T_1(x, y))\), where \(S_{nm}(x)\), is the Student distribution function or \(P_1 = 1-p\), where \(p\) is a level of ‘zero’ hypothesis significance.

### Results

The average transferrin iron concentration in the blood of animals with small and large tumors (29.6 ± 10.7 and 27.9 ± 6.7 \(\mu M\), respectively) was less than in the control group (33.8 ± 5.8 \(\mu M\), with \(P_1\) equal to 0.567 and 0.838, respectively (fig. 3). Transferrin concentrations in the blood of rats with small (68.8 ± 29.7 \(\mu M\)) and large (55.7 ± 26.2 \(\mu M\)) tumors exceeded the control level (43.9 ± 14.3 \(\mu M\), \(P_1 = 0.888\) and 0.608, respectively (fig. 3). Transferrin saturation in the blood was considerably lower than the control data (40.9 ± 13.7%), in animals with small (25.1 ± 13.7%, \(P_1 = 0.932\) and large...
(28.1 ± 9.2%, P₁ = 0.902) tumors, thereby indicating hypoperferemia (fig. 3).

Ferritin iron concentrations in the blood and liver exceeded the control level for both small and large tumors (fig. 4) and were significantly higher (P₁ = 0.99) in the livers of rats with large tumors. Stored iron content increased considerably (twofold to fourfold) compared to the control data in all brain tissues under study, in even the small tumors. The index was twofold higher than the reference value in the hypothalamus and the perifocal zone for both small and large tumors. In small tumors, a statistically significant correlation was observed between the stored iron concentration in the blood and the hypothalamus (r = 0.625; p = 0.040) as well as between the stored iron concentration in the tumor, the peritumoral tissue, and the cortex of the right brain hemisphere (p < 0.05). For large tumors a correlation was observed between stored iron concentrations in the peritumoral tissue, the cortex of the right brain hemisphere, and the hypothalamus (p < 0.002).

The chelatable iron concentration increased significantly in liver tissues and in all brain tissues except the hypothalamus (fig. 4). For small tumors, there was a statistically significant correlation (r > 0.99; p < 0.001) between the chelatable iron concentration in the liver and all brain tissues studied. For large tumors, a significant correlation was observed between chelatable iron and stored iron concentration in the blood (r = 0.744; p = 0.021) and in all brain tissues: tumor tissue (r = 0.709; p = 0.032), the peritumoral tissue (r = 0.780; p = 0.013), the cortex of the right brain hemisphere (r = 0.712; p = 0.032), and the hypothalamus (r = 0.834; p = 0.005). These correlations are indicative of the involvement of all brain tissues and the organism as a whole, in brain tumor development.

A combined statistical treatment of the differences between experimental and control data sets on non-heme iron exchange parameters is given in table 2. With small brain tumors there were 19 combinations with a normalized distance between data sets, P₁ exceeding 0.98, where-
as with large brain tumors there were 23. The high \( P_1 \) values (close to the unit) and an increase in the distance between samples upon combined parameter treatment indicate that the parameters were informative and the multifactorial approach was effective for estimating the distance between samples at glial brain tumors in rats. Normalized distance between samples as a function of the combination of non-heme iron exchange parameters could be introduced as an essential parameter in designing a risk model of glial brain tumors. In the case of animals with small tumors, combinations of indices with maximum distance between samples include transferrin saturation in the blood and the stored iron concentration in the liver and blood for both groups of animals, as well as the stored iron concentration in the tumor tissue and the cortex of the right brain hemisphere.

Non-heme iron exchange indices for the blood and biopsy samples taken from patients with glial brain tumors (table 1) show that the transferrin iron concentration values in the whole blood (19.5 ± 3.6 \( \mu M \)) were markedly and significantly higher (\( P_1 = 0.997 \)) than control data determined in healthy adults (15.7 ± 2.9 \( \mu M \)). The transferrin iron concentration in blood cells was also significantly (\( P_1 = 0.982 \)) higher in patients (8.6 ± 5.5 \( \mu M \)) than in healthy adults (5.3 ± 3.8 \( \mu M \)). The MtHb iron level in the blood of patients (1.1 ± 0.2 relative units) was threefold lower than in the group of control subjects (3.1 ± 1.0 relative units), and this difference was statistically significant (\( P_1 = 1 \)).

The ferritin iron indices increased considerably (threefold to fourfold) and significantly in the tumor tissue (135 ± 129 \( \mu M \)) and in peritumoral zone I (209 ± 160 \( \mu M \)) relative to peritumoral zone II, which was at a distance from the tumor. The ferritin iron indices also increased relative to the ferritin iron levels in patient blood and plasma (table 3), therefore, one can discriminate between normal and tumor tissue on the basis of ferritin iron con-

**Table 2.** Mehalanobis distance, \( D_I \), and normalized distance, \( P_I \), between experimental and control data sets for rats with small and large tumors as a result of the combined treatment of seven informative parameters of non-heme iron exchange: transferrin iron concentration, [Tf-Fe]bl, transferrin saturation in the blood, %Tfbl, and stored iron concentration [Ft-Fe] in the tumor tissue (t), the cortex of the right brain hemisphere (c), the hypothalamus (hy) and the liver (lv).

<table>
<thead>
<tr>
<th>Parameter combination</th>
<th>Small tumors (( \leq 3 \text{ mm}, n = 8 ))</th>
<th>Large tumors (( &gt;8 \text{ mm}, n = 6 ))</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>( D_I )</td>
<td>( P_I )</td>
</tr>
<tr>
<td>[%Tfbl, [Ft-Fe]lv]</td>
<td>14.432</td>
<td>0.998</td>
</tr>
<tr>
<td>[Tf-Fe]bl, [%Tfbl, [Ft-Fe]lv]</td>
<td>15.372</td>
<td>0.995</td>
</tr>
<tr>
<td>[%Tfbl, [Ft-Fe]t, [Ft-Fe]lv]</td>
<td>15.011</td>
<td>0.994</td>
</tr>
<tr>
<td>[%Tfbl, [Ft-Fe]c, [Ft-Fe]hy, [Ft-Fe]lv]</td>
<td>14.699</td>
<td>0.994</td>
</tr>
<tr>
<td>[%Tfbl, [Ft-Fe]bl, [Ft-Fe]t, [Ft-Fe]lv]</td>
<td>6.279</td>
<td>0.994</td>
</tr>
<tr>
<td>[%Tfbl, [Ft-Fe]hy, [Ft-Fe]lv]</td>
<td>14.438</td>
<td>0.993</td>
</tr>
<tr>
<td>[%Tfbl]c, [%Tfbl, [Ft-Fe]lv]</td>
<td>–</td>
<td>0.959</td>
</tr>
<tr>
<td>[%Tfbl]c, [Ft-Fe]lv</td>
<td>–</td>
<td>0.944</td>
</tr>
<tr>
<td>[%Tfbl]t, [%Tfbl]hy, [Ft-Fe]lv</td>
<td>–</td>
<td>0.932</td>
</tr>
<tr>
<td>[%Tfbl]t, [Ft-Fe]hy</td>
<td>–</td>
<td>0.898</td>
</tr>
<tr>
<td>[%Tfbl]t, [Ft-Fe]hy, [Ft-Fe]bl</td>
<td>–</td>
<td>0.777</td>
</tr>
<tr>
<td>[%Tfbl]t, [Ft-Fe]hy, [Ft-Fe]bl</td>
<td>–</td>
<td>0.605</td>
</tr>
<tr>
<td>[%Tfbl]t, [Ft-Fe]hy</td>
<td>–</td>
<td>0.567</td>
</tr>
</tbody>
</table>

Only 6 combinations with \( P_1 > 0.99 \) in the order of \( P_1 \) decrease are represented from a total of \( 2^7 \) possible combinations as well as the \( P_1 \) values upon separate treatment of every parameter (at the end of the table).

**Table 3.** Concentration of ferritin iron [Ft-Fe] and iron chelatable with deferoxamine B [Df-Fe] in the blood, glial tumor tissue, and peritumoral zones I and II of patients with glial brain tumors (mean ± SEM).

<table>
<thead>
<tr>
<th>Parameter</th>
<th>Blood</th>
<th>Tumor tissue</th>
<th>Peritumoral zone I</th>
<th>Peritumoral zone II</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Ft-Fe], ( \mu M/l )</td>
<td>44 ± 10</td>
<td>155 ± 39</td>
<td>203 ± 55</td>
<td>48 ± 9</td>
</tr>
<tr>
<td>( P_1 )</td>
<td>0.9939</td>
<td>0.9997 NS</td>
<td></td>
<td></td>
</tr>
<tr>
<td>[Df-Fe], ( \mu M/l )</td>
<td>5.4 ± 0.6</td>
<td>17.8 ± 2.9</td>
<td>22.2 ± 3.9</td>
<td>6.5 ± 0.1</td>
</tr>
<tr>
<td>( P_1 )</td>
<td>0.9994</td>
<td>1.000 NS</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

NS = Nonsignificant difference between data sets (\( P_1 < 0.95 \)).
The chelatable iron concentration increased significantly in the tumor tissue and in peritumoral zone I relative to peritumoral zone II, whole blood, and blood plasma. The differences we detected were statistically significant. The most informative parameter combinations ensuring the maximum normalized distance between data sets on non-heme iron exchange parameters in patients with glial brain tumors included the concentration of transferrin iron in the blood, transferrin saturation in the blood, and ferritin/hemosiderin iron concentration in the tumor tissue (table 4).

**Discussion**

Non-heme iron indices in patient groups and experimental animals are generally investigated using magnetic properties of non-heme iron species [14–19]. The phenomenon of ESR has its origin in splitting the energy levels of atomic species with nonzero electron magnetic moment in an outer, static magnetic field (electron Zeeman effect) [21]. In practice, the magnetic field is swept through the resonance while the frequency is kept constant, thus modulating the magnetic field to improve the signal-to-noise ratio. The ESR signal registered is the first derivative of an adsorption line.

Contrary to the common methods of non-heme iron status estimation, ESR method allows analysis of whole blood microsamples, achieving specificity in transferrin iron determination and allowing the determination of transferrin saturation in whole blood [14]. In addition, the new index – concentration of transferrin iron associated with blood cells – can be calculated from blood and plasma transferrin iron concentrations by taking into account the hematocrit index.

Our data show that hypoferremia was developed upon glial brain tumor formation in the rat model, whereas hypoperferremia was detected in patients with spontaneous tumor formation. These findings indicate that increased stored and chelatable iron in the tumor tissue and the adjacent peritumoral tissues are features common to glial brain tumor formation in both experimental animals and in patients.

These abnormalities can occur at any stage of iron regulation including iron uptake, storage and release, utilization, and regulation. It has been shown that non-transferrin-mediated mechanisms are involved in iron influx into brain cells [22], mediated by DMT1 coupled with ferric reductase Dcytb expressed by astrocytes [23]. Actively dividing astrocytic and tumor astroglial cells exhibit considerably higher expression of the DMT1 than nonproliferating cells [24]. The resultant increase in LMWI level leads to oxidative stress exacerbation, activation of ferritin synthesis [25] and, hence, to iron sequestration. This correlates with the up-regulation of manganese superoxide dismutase in human glioblastomas [26]. The immunosuppressive effect of ferritin [27] can, in turn, promote tumor growth.

Two stress proteins, heme oxygenase-1 (HO-1) and NO synthase, could also contribute to the increase in the redox-active iron pool and iron deposition, as reported in the present work. Induction of HO-1 [28] results in increased bilirubin and available iron and ferritin levels. Induction of the glial HO-1 was shown to lead to iron sequestration in mitochondria and to mitochondrial insufficiency in human central nervous system disorders [29]. Both hypoxia and nitrogen monoxide can also induce ferritin synthesis [30].

Stored iron accumulation during glial brain neoplasm formation could explain the robust association between glioma and epilepsy [31], as epilepsy could be the initial
and only clinical manifestation of a glioma [32, 33]. Iron deposition is a prominent feature of human post-traumatic epilepsy [34]. Subcranial injection of iron-containing products into the cortex was shown to produce a chronic epileptic focus in experimental models [35]. The efficacy of antioxidants in preventing iron-induced epileptiform activity in animal models suggests an important role for oxidative injury in the development of experimental epilepsy [36]. We have also found plausible shifts in iron status in animals using the Kindling epilepsy model. In particular, considerable increases in the ferritin iron level in hippocampal tissue were reliably established.

MtHb is thought to be a marker for oxidative stress. Surprisingly, in patients with glial brain tumors we found a statistically significant threefold decrease in MtHb levels in the blood compared to reference data. One could speculate that the decrease in steady-state MtHb iron concentration in the blood occurred due to the release of hemin, which, in turn, can contribute significantly to increased proliferation of cancerous cells, as was shown in vitro for human glioma cell lines [37]. Alternatively, adaptive activation of the NADH-methemoglobin reductase could account for the decreased MtHb in tumor patients [38].

We found that the most informative parameter combinations distinguishing between patients and control subjects included transferrin iron concentration in the blood, transferrin saturation in the blood, and ferritin/hemosiderin iron concentration in the tumor tissue.

Iron accumulation in tumor tissue is associated with an increase in relative intensity of ESR components with magnetic anisotropy (fig. 2b, c), which can be attributed to the magnetic susceptibility anisotropy of ferritin/hemosiderin aggregates in tissues. The observed ESR shift on sample rotation in the spectrometer resonator is indicative of the local magnetic fields in aggregates of the order of at least several millitesla. One could speculate about possible effects of the magnetic fields due to ferritin/hemosiderin iron in pathological development, in particular, accounting for the catalysis of free radical oxidation in steady-state magnetic fields of 10–80 mT [39], or in the in vitro inhibition of phagocytic activity of immunocompetent cells placed in applied static magnetic fields of 25–150 mT [40].

The efficiency of iron chelators in the treatment of neuroblastoma, which is also characterized by increased ferritin and hemosiderin levels, has been observed both in cell culture and in patients [41, 42]. A set of iron chelators has already been shown to inhibit tumor cell growth and to have significant in vivo antineoplastic activity. It has been demonstrated that some iron chelators can penetrate the blood-brain barrier and that desferrioxamine is effective in diminishing central nervous system iron accumulation [9, 43]. In this context it may be worthwhile to assess chelators for treatment of glial brain tumors.

**Conclusion**

In the present study we found that patients with malignant glial brain tumors exhibited hyperferremia in the blood, whereas experimental glial brain tumors in rats were marked by hypoferremia. Tumor tissue and adjacent peritumoral tissue were considerably enriched in stored and chelatable iron upon glial brain tumor formation in both the rat experimental model and in humans.

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References


