SHORT REPORT

Spontaneous intralesional haemorrhage in dysembryoplastic neuroepithelial tumours: a series of five cases


Abstract

Five patients with dysembryoplastic neuroepithelial tumour (DNT) showing extensive secondary haemorrhage, a finding not previously associated with these neoplasms, are described. The clinical presentations, neuroimaging findings, and histopathological features of these patients are reviewed. One patient, a previously asymptomatic 12 year old girl, presented with an acute intracerebral haemorrhage into a DNT. A further four young adults with histories of intractable partial and generalised seizures dating from childhood showed significant chronic haemorrhages within DNT, the MRI appearances in one patient giving a false impression of a cavernoma. Histopathology disclosed vascular abnormalities within these tumours which, together with other factors discussed, may have predisposed these tumours to haemorrhage.

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Keywords: DNT; epilepsy; hemorrhage; MRI

Dysembryoplastic neuroepithelial tumour (DNT) is a low grade, mixed neuronal and glial tumour usually associated with medially intractable complex partial or generalised seizures which date from childhood. The prognosis of these neoplasms is usually excellent after their identification with MRI and subsequent surgical excision and there has been only one report of malignant transformation.

It is well recognised that an acute intracerebral haemorrhage may mark the initial presentation of a primary intracranial glial tumour of both high and low grade and it has been suggested to be a particular liability in oligodendrogliomas, related to the prominent arculate vascular patterns of these tumours. Vascular patterns similar to oligodendrogliomas are often present in DNT but there has previously been only one report documenting an asymptomatic haemorrhage within a DNT.

We present five further DNTs with spontaneous haemorrhage, which complicated either the clinical presentations or the pathological diagnoses.

Methods

The five cases of DNT presented were encountered in the neurosurgical material examined at the National Hospital over a 2 year period (1994–6); histological sections from patient 1 were initially referred from the Morriston Hospital, Swansea, for review by one of us (TR). A review of all other DNTs on the hospital surgical pathology data base (28 tumours) found no similar cases. The clinical data from all patients were reviewed including seizure history, neuroimaging features, operative procedures, and follow up. The resected tissue was formalin fixed, paraffin embedded, and reviewed using conventional stains, including Perl’s techniques for iron and immunohistochemistry (GFAP 1:400 Dako, and synaptophysin 1:20 Dako) for confirmation of the glial and neuronal components of the tumour. In three of the cases, intraoperative cytological smear preparations stained with toluidine blue were also available for review (patients 2, 4, and 5).

CLINICAL AND NEURORADIOLOGICAL FINDINGS AND SURGICAL TREATMENTS

Case 1

A previously well 12 year old girl presented with a short history of headache and confusion. Brain CT performed on admission showed a large acute intracerebral haemorrhage in the right temporoparietal region and angiography was negative. Six months later an MRI was performed for persisting headache which disclosed a well defined cystic mass in the right temporoparietal region and angiography was negative. Six months later an MRI was performed for persisting headache which disclosed a well defined cystic mass in the right temporoparietal region and angiography was negative. Six months later an MRI was performed for persisting headache which disclosed a well defined cystic mass in the right temporoparietal region and angiography was negative. Six months later an MRI was performed for persisting headache which disclosed a well defined cystic mass in the right temporoparietal region and angiography was negative. Six months later an MRI was performed for persisting headache which disclosed a well defined cystic mass in the right temporoparietal region and angiography was negative.
developed a left homonymous superior quadrantanopia but otherwise made a full recovery, is well at 1 year follow up, and has not at any time had seizures.

**Cases 2–5**

The clinical and operative details of these cases are summarised in the table. All patients had medically refractive seizures dating from childhood but none had histories of acute neurological deterioration to suggest episodes of intratumorous haemorrhage. The MRI appearances showed well demarcated, nodular, and cystic intracortical lesions involving temporal or frontal lobes. The signal characteristics in patients 3, 4, and 5 were typical of a DNT with lesional hypointensity on T1 and hyperintensity on proton density and T2 weighted images. However, in patient 2 the MRI was not typical, showing central CSF-like signal intensity, a peripheral rim of hypointensity on proton density, and T2 weighted images suggesting previous haemorrhage (figure C) and surrounding hyperintensity in keeping with gliosis. These
features led to a preoperative radiological diagnosis of a cavernoma. Preoperative serial MRI in patient 5 confirmed progressive enlargement of the tumour with cystic change between 1991 to 1995. There was no radiological evidence of calcification either on CT or with MR in any of these tumours and in patients 2 and 3 contrast administration showed no enhancement and angiograms were negative.

**HISTOLOGICAL FINDINGS**

In two of three tumours in which cytological smear preparations were available for review, proliferations of astrocytes containing abundant haemosiderin pigment were present suggesting a cellular reaction to an old haemorrhage (patients 2 and 4); the third (patient 5) showed features of a low grade glioma consistent with a DNT. Surgical resections included two total (patients 2 and 3) and three partial (patients 1, 4, and 5) lesionectomy specimens, and histological examination of these confirmed intracortical neuronal-glial tumours with a complex amalgam of architectural patterns as described in DNT. In some areas the normal cortex was replaced by tissue of a loosely structured mucinous nature, in which randomly distributed mature appearing nerve cells were “suspended” (figure D) and smaller oligodendroglial-like cells were clustered around thin walled capillaries representing the specific glioneuronal element of DNT. Well defined tumour nodules composed of rounded oligodendroglial-like cells and intermingled mature nerve cells were seen and also areas with pilocytic astrocytoma-like growth pattern were present in two (patients 1 and 2). All patients showed extension of tumour into the leptomeninges. Formation of cystic cavities of varying size were present within all the

<table>
<thead>
<tr>
<th>Patient Sex</th>
<th>Age at onset of seizures (y)</th>
<th>Seizure type and approximate frequency before surgery</th>
<th>Age at surgery (y)</th>
<th>Tumour location and surgical procedure</th>
<th>Outcome and duration of follow up</th>
</tr>
</thead>
<tbody>
<tr>
<td>2 Male</td>
<td>7</td>
<td>Partial (1/week) and generalised seizures (1/week)</td>
<td>24</td>
<td>Right frontal. Removal of cystic mass (total excision).</td>
<td>Only one seizure since surgery. Postoperative EEG shows no epileptiform activity. 2 year follow up Seizure free. 18 month follow up.</td>
</tr>
<tr>
<td>3 Male</td>
<td>9</td>
<td>Complex partial seizures and generalised seizures (2–3/week), Absence and generalised seizures (2/week),</td>
<td>27</td>
<td>Left temporal. Lesionectomy; lower middle and inferior temporal gyrus.</td>
<td>Continues to have seizures, although decreased in frequency (2/month), 18 month follow up. Marked reduction in seizure frequency. 1 year follow up.</td>
</tr>
<tr>
<td>4 Female</td>
<td>6</td>
<td>Complex partial seizures (5–10/month) and generalised seizures (5/year),</td>
<td>20</td>
<td>Left frontal. Subtotal (80%) excision: due to tumour location in motor cortex.</td>
<td></td>
</tr>
<tr>
<td>5 Female</td>
<td>6</td>
<td>Right frontal. Cystic lesion extending to cortex. Cyst drained and partial resection of tumour.</td>
<td>41</td>
<td></td>
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</table>
tumours, many close to the meningeal surface (figure E). Adjacent cortical dysplasia was noted in three patients (1, 2, and 3).

Extensive intratumorous haemosiderin deposition was confirmed with Perl’s staining; the iron pigment was abundant in cyst walls and also diffusely impregnating tumour cells (figure F). The tumour vasculature was composed predominantly of delicate thin walled branching capillaries, typical of the arcuate pattern previously described in DNT, and was present in both tumour nodules and the glioneuronal elements of all cases. These vessels traversed through the mucinous matrix of the tumour where they appeared relatively unsupported by astrocytic processes. Such vessels may be more vulnerable to mechanical shearing during episodes of minor cerebral trauma, a not uncommon occurrence in patients with repeated and poorly controlled generalised seizures. Also present in two patients were glomeruloid-like tufts of proliferating capillaries were present in close proximity to areas of haemorrhage. Extensive mineralisation of vessels (figure G) was found in two tumours (patients 1 and 3). In none of the cases were vascular endothelial proliferations characteristic of malignant astrocytoma seen and there was no evidence of vascular thrombosis, amyloid deposition, or angiitis.

Discussion
Spontaneous intratumorous haemorrhage may mark the initial presentation of both low and high grade glial neoplasms. DNTs are distinctive low grade glial-neuronal neoplasms typically associated with intractable epilepsy dating from childhood. There is a single, previous report of a DNT in which a chronic haemorrhage was found but an acute presentation with spontaneous intratumorous haemorrhage in a histologically established DNT has not been previously documented. In a further four DNTs we have histological evidence of chronic haemorrhage which in the absence of episodes of acute neurological deterioration, we can suspect has accumulated over time from small clinically silent bleeds. In three of the four patients, the haemorrhage was not visualised by MRI, which may indicate smaller amounts of haemosiderin deposition in these patients.

None of the patients had evidence of a coagulopathy and we examined the tumour vasculature as a potential factor which may have predisposed these tumours to haemorrhage. The angiograms, when performed, were normal and enhancement with contrast MRI was only seen focally in patient 1, 6 months after an acute cerebral haemorrhage. Although this enhancement may have been interpreted as neovascularisation of an organising haematoma within the tumour, it was considered more likely to represent abnormal vessels within the tumour itself. Enhancement on MRI within DNTs is not in itself an unusual finding and has been reported in up to 36% of patients in one series and in six of 16 patients in another.

Histologically, the predominant vascular patterns in the neoplasms were fine delicate, branching capillary networks, also termed arcuate or retiform vessels. Similar vascular patterns have been shown to be significantly associated with tumour haemorrhage in a study of gliomas. It was also noted in this same study that haemorrhage within oligodendrogliomas correlated with calcification of such tumour vessels. We also found dystrophic calcification or mineralisation of small vessels in two of the present cases. It is not inconceivable that such calcified vessels may have impaired haemostatic vasoconstrictive responses and so contribute to the extent of haemorrhage. Additionally, we noted that the fine capillaries within the DNT characteristically traverse through the abundant mucoid matrix of the tumour, relatively unsupported by astrocytic processes. Such vessels may be more vulnerable to mechanical shearing during episodes of minor cerebral trauma, a not uncommon occurrence in patients with repeated and poorly controlled generalised seizures. Also present in two patients were glomeruloid-like tufts of proliferating capillaries in close proximity to areas of haemorrhage suggesting a causal link, although in a previous study of gliomas no significant association between the presence of haemorrhage and glomeruloid vascular pattern was found. These vascular abnormalities have occasionally been noted in previous published series of DNTs; however, in the only previously documented case of a DNT with haemorrhage, no mention of abnormal vasculature was recorded.

Cystic degeneration, a not infrequent finding in DNT, was present in all the cases on MRI and confirmed histologically, many of the cysts containing old blood pigment suggesting they may have been a factor contributing to the accumulative size of the haemorrhages. Also, invasion of the leptomeninges by these superficial tumours was common to all, with large cysts closely opposed to the frontal or temporal cortical surface (figure C) which may also have increased their vulnerability to traumatic haemorrhage.

We can hypothesise that the iron deposited within these tumours played a functional role in seizure propagation and worsening of the epilepsy in these cases. Iron is known to precipitate epilepsy both in clinical and experimental situations; free radical formation and lipid peroxidation of neural membranes and excitatory neurotransmitter release are possible mechanisms for seizuregenesis in this model of epilepsy. The role of iron in enhancing epileptiform activity within these DNTs, however, remains highly speculative, particularly as in patient 1 there was no history of seizures either before or after partial tumour resection.

The are several diagnostic considerations arising from this group of patients: firstly, that DNT may rarely present with an acute intracerebral haemorrhage as in the first patient of this series. This case also highlights the potential value of MR neuroimaging in young patients with angiogram-negative intracranial haemorrhage as has been noted in a previous study. It was also remarkable in the absence of a history of seizures which although unusual for DNT, has occasionally been seen. We have also shown that clinically silent, chronic haemorrhages may accumulate in DNT, and significantly alter the MRI appearance giving a false impression of a cavernoma. Lastly, the interpretation of the
intraoperative cytological smear preparations may be complicated by the old haemorrhage and reactive gliosis present in such DNTs. We therefore conclude that the presence of haemosiderin pigment on smears which otherwise show appearances typical of a DNT should not deter the pathologist from making this diagnosis and thus reassure the surgeon to proceed with the appropriate lesionectomy.

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