Abstract

Aims—To study the mechanism of action of steroids in patients with peritumorous oedema.

Methods—To investigate early cerebral metabolic changes proton magnetic resonance spectroscopy (1H-MRS) was used before and 11 to 14 hours after treatment with dexamethasone (12 mg oral loading and 4 mg four times daily maintenance).

Nine patients (two men, seven women, mean age 54) with pronounced oedema associated with various intracranial tumours (two astrocytomas, three meningiomas, two glioblastoma, and two metastases) were examined using MRI and MRS. SE1500/135 volume selected MRS (mean volume 21 ml) were performed on an oedematous region and a contralateral region. All spectra were acquired with and without water suppression. Metabolite peak area ratios were determined.

Results—Regions of oedema had significantly (P < 0.01) higher unsuppressed water than the contralateral regions, as expected. There was no change at this early time point after dexamethasone. The ratio of the area of choline containing compounds to that creatine and phosphocreatine compounds was determined after which the serial ratios of these before and after were calculated (a serial ratio of 1.0 would indicate no change in the choline to creatine ratios after steroid administration). The mean serial ratios for the area of oedema were 1.02 (SEM 0.08) and 1.10 (0.08) for the contralateral volume of interest, indicating no significant changes. However, significant changes (P < 0.02) were found in the N-acetyl-aspartate (NAA)/choline serial ratios (0.86 (0.06) in the area of oedema, 1.20 (0.10) in contralateral brain) and the NAA/creatinine serial ratios (0.86 (0.08) for the oedema, 1.25 (0.11) in contralateral brain).

Conclusions—Such rapid changes may be explained either by relatively large alterations in the relaxation characteristics of NAA or, more controversially, by actual changes in the amounts of NAA. It is proposed that steroids act primarily by causing early metabolic changes that are later expressed in improvements in intracranial volume relations.

Keywords: brain; dexamethasone; peritumorous oedema; spectroscopy; metabolism

Steroids are routinely employed in the treatment of patients with tumours of the CNS yet their mode of action remains obscure. In particular, the rapid and dramatic clinical improvement seen in some patients is difficult to explain on the basis of changes in blood-brain barrier permeability, blood flow or volume, decrease in CSF production, or resolution of peritumorous oedema. We have utilised proton magnetic resonance spectroscopy (1H-MRS) to study peritumorous oedema before and after treatment with dexamethasone to examine possible early metabolic changes.

Materials and methods

Nine patients (two men, seven women, mean age 54) with primary space occupying brain tumours were examined using a 1.5T Siemens Magnetom with a standard quadrature transmit/receive head coil. Collimated light localisers were used to serially reposition the patient’s head with respect to the coil and magnet. SE1500/135 (256 averages) volume selected MRS was performed on the oedematous region and a contralateral region. All spectra were obtained with water suppression (256 averages) and without it (one average). The chemical shift selective saturation CHESS sequence was used for water suppression. Global and then local shimming were used to bring the unsuppressed water linewidth (full width at half maximum) in the selected volume of interest (VOI) to within 10 Hz.

A standard T2 weighted MRI sequence (SE2200/80, 192 × 256, 20 × 5 mm slices in the axial orientation) was acquired both before and after treatment to ensure that no macroscopic changes in brain water had occurred over this short time frame. In addition “scout” sets of sagittal and coronal images were acquired using an SE400/15 (128 × 256, 10 × 10 mm slices) to guide positioning of the VOIs in conjunction with the T2 weighted axial image (fig 1). These were placed within the oedema but excluding the tumour itself. To optimise signal to noise ratios VOIs were chosen to encompass as much oedematous white matter as possible, but not to be near the inner table of the skull to avoid line broadening and lipid contamination of the spectra by marrow and scalp. A second VOI was placed...
Early changes in peritumorous oedema and contralateral white matter after dexamethasone: a study using proton magnetic resonance spectroscopy

Figure 1 Images showing typical positioning of VOI over oedema: (A) axial, (B) sagittal, (C) coronal.

Figure 2 T2 axial slice showing typical positioning of contralateral VOI.

The interactive phase correction of the frequency data to make the baseline as linear as possible over the frequency range of the metabolites of interest. There is always some degree of subjectivity inherent in choosing integration limits for peak area calculations so to avoid introducing between observer errors all peak area calculations were performed by one observer (BC).

Ethical approval was obtained for this study and each patient signed a consent form before participating. Recruitment of patients was diffi-
cult due to the fact that referring physicians had often commenced steroid treatment before transfer. In addition, ethical considerations in delaying treatment even for the few hours necessary to secure an MR time slot meant that patients with florid symptoms and signs of raised intracranial pressure were not studied. None of the patients showed dramatic improvement after dexamethasone over the time scale of the study.

Results

Comparison of the T2 weighted images disclosed no visible changes in oedema contrast or extent after dexamethasone treatment. Similarly there were no significant serial changes in the unsuppressed water peaks either in the oedematous or normal sides, which support previous findings. The VOIs over oedema had significantly higher (P < 0.01) unsuppressed water peak areas than those over the contralateral side, as expected.

The area ratios calculated were N-acetyl aspartate (NAA)/choline (Cho), NAA/creatine + phosphocreatine (Cr) and Cho/Cr, where Cho refers to the peak reflecting choline containing compounds and Cr refers to the peak reflecting creatine and phosphocreatine compounds. As expected the VOI over oedematous brain before dexamethasone treatment showed some reduction in the NAA/Cho ratios with a mean (SD) ratio of 1.52 (0.89) compared with 1.84 (0.24) in the contralateral VOI. For NAA/Cr ratios the values were 1.78 (0.36) and 1.90 (0.28) respectively. Choline to creatine ratios differed more in the oedematous VOI with a mean of 1.79 (1.45) compared with 1.04 (0.19) on the contralateral side.

We are, however, interested specifically in the changes to these relative peak areas after drug administration. Thus for both the oedematous side and the contralateral side we calculated the following serial ratios: post-NAA/Cho/pre-NAA/Cho; post-NAA/Cr/pre-NAA/Cr; post-Cho/Cr/pre-Cho/Cr. If there had been no change then all these serial ratios would yield a value of 1.00. Table 1

<table>
<thead>
<tr>
<th>Serial ratios</th>
<th>Oedema</th>
<th>Contralateral side</th>
<th>P value</th>
</tr>
</thead>
<tbody>
<tr>
<td>NAA/Cho</td>
<td>0.86 (0.16)</td>
<td>1.20 (0.10)</td>
<td>&lt; 0.02</td>
</tr>
<tr>
<td>NAA/Cr</td>
<td>0.86 (0.08)</td>
<td>1.25 (0.11)</td>
<td>&lt; 0.02</td>
</tr>
<tr>
<td>Cho/Cr</td>
<td>1.02 (0.08)</td>
<td>1.10 (0.08)</td>
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</tr>
</tbody>
</table>

Mean 1.00 would represent no change.

Figure 3 Mean (SEM) serial peak area ratios for oedema and contralateral side.

Figure 4 Examples of the small but observable changes in spectra after dexamethasone for (A) oedematous and (B) contralateral VOI, with relative reductions in NAA in the oedematous region, and relative increases in the contralateral region. The predexamethasone spectra have been displaced vertically for ease of comparison.
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Negative peaks at the chemical shift position corresponding to lactate were found in all patients in the oedema spectra, and in one instance in the contralateral spectra. These peak areas were generally small compared with the metabolite peak areas and showed no evidence of change after dexamethasone.

Discussion
Steroids have been used in the treatment of brain tumours since the early 1950s when Ingraham et al noted benefit from their use in patients with craniopharyngioma. However, it was not until the report of Galich et al in 1961 that the wider application of steroids for all CNS tumours was recognised. Over the past 30 years, numerous studies have been performed to attempt to elucidate the mechanisms of action but none have been able to explain the rapid clinical response. In the present study, significant metabolic changes were seen within 14 hours, both in the zone of oedema around the tumour and in the contralateral hemisphere. A finding all the more surprising considering the largely asymptomatic nature of many of the patients and the lack of obvious clinical change. We propose that steroids have a primary metabolic action—of which the changes in NAA seen in this study represent only a small part. Any changes in blood-brain barrier permeability and resolution of the oedema being secondary to the altered metabolic milieu. The time scale of these changes is in keeping with possible enzyme induction or inhibition.

PROTON MAGNETIC RESONANCE SPECTROSCOPY
Proton magnetic resonance spectroscopy (H-MRS) is being used increasingly often as a non-invasive method for analysing regional cerebral metabolism in vivo. Changes in H-MRS have been shown in multiple sclerosis, epilepsy, cerebral infarction, AIDS, and brain tumours. The hope has been to develop a tumour specific metabolic profile to act as a fingerprint and aid in preoperative diagnosis. After, initially, a lack of reliable correlation, Preul et al have reported for certain tumours a 99% correlation between spectroscopic data and final pathological diagnosis. As a general rule in cerebral tumours, there is a decrease in the relative concentration of NAA and creatine/phosphocreatine together with an increase
in choline and lactate.22 Despite the fact that dexamethasone is used routinely in these patients no previous study has analysed the changes in the spectra as a consequence of this drug. 

Forty years after its identification by Tallan et al, the role of NAA in cerebral metabolism remains uncertain.23 It is found only in the nervous system and is second only to glutamate in total concentration of free amino acids.24 In 1H-MRS, NAA produces a sharp spectral peak suggesting that it may be mobile and in free solution.24 Although regarded widely as a neuronal marker, there is still debate about its site of production. In particular, apparent increases in NAA seen during the recovery phase of various pathological states casts some doubt on this assumption.25-27

Some have suggested that NAA is involved in lipid synthesis in the production of myelin;28-30 or as a regulator of protein synthesis; or as a storage form of aspartate; or it is a breakdown product of another compound.24

In the present study we focused on changes in the zone around the tumour and found that the relative concentration of NAA apparently either decreased on the side of the oedema or increased on the contralateral side within 14 hours after treatment with dexamethasone. This implies that either the amount of NAA is changing, or there is a change in the relaxation behaviour of NAA. The degree of changes in relaxation time necessary to produce such large changes in apparent amounts of NAA present can be estimated by applying the standard spin echo signal equations with these pulse timing variables, and using previous estimates of NAA relaxation times26-27 of 1650 ms (T1) and 330 ms (T2). Such calculations show that to account for the increased NAA signal in the contralateral region, either a 40% reduction in T1 or a 170% increase in T2 would be required. The apparent 14% reduction in oedematous NAA signal would require around a 30% increase in T1 or a 30% decrease in T2.

Such large T2 changes in oedematous NAA are not without precedent (although over the much longer time course of oedema resolution). Using two different TEs and two different TRs Kamada et al were able to estimate T1 and T2 for the oedematous NAA associated with brain tumours.26-27 The T1 showed similar values to the NAA of normal brain but the T2 was found to have decreased by about 50%.26-27 The use of steroid treatment is not mentioned, but it is likely that all the patients had received treatment so their results would relate to our findings after treatment. In two patients who had long term follow up a gradual “normalisation of T2” occurred as the oedema resolved.26-27

No changes in contralateral NAA were found.

Large changes in metabolite T2 values over a short time after initiation of treatment would therefore be required to explain these results. Changes in binding (changes in relaxation time) may indeed provide the explanation but it is also possible that we have detected increases in the transport out of a free and mobile molecule because of metabolic improvement. Other possibilities include increased utilisation (perhaps for lipid or protein synthesis for repair of damaged myelin) or increased breakdown (white matter having three times as much hydrolyase activity as grey matter—despite only having half the concentration of NAA).24

If we are seeing real changes in NAA rather than relaxation effects, the apparent rise in NAA on the contralateral side is more difficult to explain but could represent either increased production or decreased utilisation or breakdown of this compound. McIntosh and Cooper have previously reported that drugs that normally raise cerebral concentrations of 5-hydroxytryptamine also increase concentrations of NAA.30

CEREBRAL METABOLIC CHANGES AFTER DEXAMETHASONE

Corticosteroids cause systemic metabolic changes which include hyperglycaemia and catabolic effects that are usually mediated via intracellular receptors acting on gene expression.31 Likewise within the brain, steroids have been shown to act both genomically and non-genomically,31 in the second case via direct action on cell surfaces, alteration in ion permeability, and by the release of neurohormones and neurotransmitters.32 In the brain, two types of receptors bind glucocorticoids with high affinity: the type I mineralocorticoid receptor and the type II glucocorticoid receptors32 and although the anatomical distribution of these receptors varies, studies have shown uptake in neurons and all classes of astrocytes (including oligodendrocytes).33 Studies using [1H]dexamethasone have shown uptake in the cerebral hemispheres of about 50% (with the remainder evenly divided between brainstem and cerebellum) in control animals and this increases to 75-80% after cerebral damage. Before trauma, 75% of the dexamethasone is located in astrocytes and 25%,35,36 neurons whereas after cerebral insult this changes to 48% bound to astrocytes and 42% to neurons.35 At the subcellular level, accumulation occurs in the microsomal, lysosomal, and cytoplasmic fractions of the damaged cells.35

Whatever the mechanism of action at the cellular level, steroids have been shown to preserve cerebral energy reserves.34-36 In particular, in the neonatal period hydrocortisone has been shown to increase brain glucose, glycogen, B-hydroxybutyrate, and ATP concentrations after decapitation in mice.34 35 Recent studies using pretreatment with small doses of dexamethasone in a neonatal rat model of hypoxia-ischaemia have shown virtually complete protection.37 Even in adult models of ischaemia—in which no beneficial effect from treatment with steroids has been shown and there is even some evidence of a detrimental effect—prior treatment with dexamethasone has maintained ATP concentrations and electrolyte balance.22,36 Both of these may indicate a preference for non-oxidative metabolism with structural and kinetic changes in rate limiting enzymes and a reversion to fetal
type enzymes. It may be, therefore, that dexamethasone acts by inducing changes in these enzymes, which in turn increase local energy reserves and improvement in cellular homeostasis.

Conclusion

The study confirms that there are significant early changes in cerebral metabolites after treatment with dexamethasone and that these changes can be monitored in vivo over the time that clinical benefit occurs. Our work also shows that the consequences of steroid administration must now be taken into account in the previous reports in which brain tumours were studied with MRS. Further studies are underway to examine changes in tissue energy status by using phosphorus MRS and to examine metabolite changes within the tumours.

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