EDITORIAL COMMENTARY

Herpes simplex virus encephalitis

Corticosteroids in herpes simplex virus encephalitis

H Openshaw, E M Cantin

The effectiveness of corticosteroids in herpes simplex virus encephalitis is not proven

Despite receiving standard antiviral therapy, an appreciable number of patients with herpes simplex virus encephalitis (HSVE) experience poor acute outcome or delayed neurological progression. It is uncertain whether all poor outcomes are due to viral cytopathology or whether an immune mediated pathogenesis also occurs. In support of an immunopathological cause, Kamei et al (see pages 1544–9 of this issue) report that corticosteroid administration was a significant predictor of favourable outcome at 3 months after HSVE infection.

Immune cells persist and elaborate cytokines in the nervous system long after the virus has entered latency. It is likely that there is ongoing antigenic stimulation, probably from low level HSV reactivation. To determine the effect of long term silencing of HSV, a clinical trial sponsored by the National Institutes of Allergy and Infectious Diseases has been designed and compares 3 months of oral valaciclovir to placebo in HSVE subjects who have completed their intravenous acyclovir course (http://clinicaltrials.gov). This long term treatment may significantly influence the CNS inflammatory cell infiltrate. Evidence suggesting that CNS inflammation has a detrimental effect comes from an experimental study in knockout mice lacking the Toll-like receptor 2, one of the receptors that mediate the inflammatory cytokine response to HSV. Despite similar CNS viral titres, lethal encephalitis occurred in a lower percentage of the knockout mice. A similar beneficial effect from corticosteroids, presumably through an anti-inflammatory mechanism, was reported in a brain magnetic resonance imaging (MRI) study of limbic HSVE in mice. Significantly less brain T2 hyperintensity was seen in animals receiving acyclovir and methylprednisolone compared to acyclovir alone at 2 and 6 months after HSV inoculation. Unlike the report of Kamei et al, where patients received corticosteroids concurrently with acyclovir, initiation of methylprednisolone was delayed until after completion of a 2 week course of acyclovir, and methylprednisolone was given only for 1 week. In the same animal model, acyclovir started 1 day after HSV inoculation reduced levels of brain chemokine mRNA expression, particularly CCL5, and concurrently administered methylprednisolone resulted in an additional decrease. If a similar reduction in chemokines occurs in patients, the extent of chronic inflammation and neuronal damage caused by HSVE would be expected to be less.

There are, of course, problems applying results from animal models to HSVE in patients. Furthermore, concerns that early corticosteroid administration may increase viral CNS spread have led many practitioners to restrict corticosteroids to patients with significant brain oedema. The retrospective study of Kamei et al should not be taken as evidence of the effectiveness of early corticosteroid treatment in HSVE. After all, the decision to use corticosteroids (as well as the preparation and schedule) was at the discretion of the treating physician. Perhaps the real predictor of outcome is not corticosteroids per se but rather some (undefined) characteristic of the infection that leads to corticosteroid use. The effectiveness of corticosteroids in HSVE can only be determined by prospective, randomised studies. Before such studies are designed, it would be helpful to have additional animal model results describing the optimum timing of corticosteroid administration as regards the effect on the CNS inflammatory cell infiltrate.

REFERENCES


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Tubercul cent meningitis

**BCG vaccination and severity of childhood tubercul cent meningitis**

**V Rajeshkehar**

BCG vaccination may lessen the severity of, rather than completely prevent, serious forms of tuberculosis

In this issue of the journal (see pages 1550–4), Kumar et al report on the efficacy of bacillus Calmette-Guerin (BCG) vaccination in favourably altering the course and outcome of tubercul cent meningitis (TBM) in children. The value of BCG vaccination for preventing tuberculosis has been debated since its introduction nearly eight decades ago. Several trials, cohort studies, and case control studies have failed to provide a definitive answer to this contentious issue. A recent meta-analysis revealed that the protective effect of BCG vaccination in infants was only around 50% against all forms of tuberculosis.1 It has been suggested that although BCG vaccination is not highly effective in preventing the illness, it does play a role in reducing the severity of the more serious forms of the disease such as TBM.2 The authors of the present report set out to study this premise and also to document the presentation of the disease in vaccinated children. The strength of their work lies in the prospective collection of their data. The authors’ data suggest that children who had been vaccinated had a milder form of the disease and consequently had a better short term outcome than those who had not received vaccination. Vaccinated children also had fewer focal neurological deficits and higher cerebrospinal fluid (CSF) cell counts. The authors speculate that vaccinated children are probably capable of mounting a better immunological response than unvaccinated children (reflected in the higher CSF cell counts) and that this is responsible for the better outcome. It is, however, unclear from the authors’ data whether there were factors other than vaccination that contributed to the severity of the illness in the unvaccinated children. It is pertinent to note that the duration of the disease at presentation to the hospital in unvaccinated children was on average 10 days longer than that in vaccinated children. 

These deficiencies apart, this study adds to the body of literature that indicates that the utility of BCG vaccination lies more in influencing the severity of the serious forms of the disease, such as TBM, than in providing absolute protection against the disease. In doing so, it supports the continuation of BCG vaccination as part of the Universal Immunization Programme (UIP) in endemic countries such as India.

**REFERENCES**


Can MRI distinguish injurious from innocuous trigeminal neurovascular contact?

**W P Cheshire**

It may be better not to operate on patients with chronic facial pain

Traditional wisdom teaches that “good surgeons know how to operate, better ones when to operate, and the best when not to operate”1.2 When treating patients with chronic facial pain, prudent application of this aphorism draws not only from experience but also from rigorous scientific investigation. Lang et al (pages 1506–9 of this issue) have strengthened the base of scientific evidence that informs the clinical decision not to recommend microvascular decompression for persistent idiopathic facial pain (PIFP).
inhibition is inadequate. Lancinating or electrical pain strikes fleetingly and repetitively in response to normally non-painful afferent stimuli, such as oral movement or light touch at a remote facial trigger zone.

But then, not all facial pains behave as trigeminal neuralgia, and craniotomy is not without risks. Although no controlled clinical trial has shown benefit from microvascular decompression in atypical, non-paroxysmal trigeminal distribution pain, microvascular decompression is occasionally tried for the individual patient with terrible intractable unexplained pain. Severely affected patients consult many physicians and may be willing to undergo invasive procedures of uncertain benefit. In such cases, magnetic resonance imaging (MRI) may suggest a rationale for surgical intervention should it happen to disclose a vessel coursing alongside the trigeminal nerve root. Recent advances in MRI spatial resolution have brought into view more of these neurovascular liaisons. Finer imaging detail, however, alone cannot distinguish the pathological from the incidental. Systematic clinical correlations are required for the sake of diagnostic clarity and treatment validity.

Lang et al utilised highly sophisticated MRI with 3D reconstruction to assess 12 subjects with PIFP. Radiologists blinded to the laterality of the pain evaluated the images for neurovascular contact, which was frequent, occurring in nine subjects. Of distinct interest is the fact that the presence of neurovascular contact did not differ between the symptomatic and asymptomatic sides, discounting any causal relationship to pain. None of the PIFP patients had morphologies of grooving, distortion, or deviation of the trigeminal root, which are considered more specific for trigeminal neuralgia.

Visualisation of neurovascular contact, it must be concluded, is a non-specific finding that should not itself be used as a convenient way of establishing a diagnosis or opting for surgery in the patient with PIFP. Satisfactory outcomes for patients are still best guided by clinical criteria, in particular, the temporal pattern, pain characteristics, and triggering factors.


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