Isotope encephalography in the management of acute herpesvirus encephalitis

A. S. BLIGH, C. M. WEAVER, AND C. E. C. WELLS

From the Departments of Radiology, Child Health, and Neurology, Cardiff Royal Infirmary, Cardiff, Wales

SUMMARY Two patients with Herpesvirus hominis (herpes simplex) encephalitis were investigated by serial isotope encephalograms using technetium-99m pertechnetate. In the first case the diagnosis was made by brain biopsy, and by successful tissue culture, and was confirmed by a significant rise in antibody titre, but in the second the laboratory evidence was exclusively serological. Necrotizing encephalitis was presumed in both cases because the illness was grave and focal signs developed in conjunction with radiological and electroencephalographic evidence of circumscribed lesions of the hemisphere. The emergence of new lesions in the brain scan at a time of clinical remission was found to be a warning of impending relapse. Such lesions had to be distinguished from areas of increasing uptake of isotope due only to local change in vascular permeability. Decay of EEG activity together with clinical signs of focal abnormality over an area of isotope concentration indicated a necrotizing lesion. Cytosine arabinoside (cytarabine), an alternative to idoxuridine as an antiviral drug, was used in both cases. A third patient, suspected in life of having a degenerative encephalopathy, was found at necropsy to have necrotizing encephalitis. Herpes infection though probable was not confirmed. The most severe lesions were in the frontal and temporal lobes, which had shown increased uptake of technetium in the encephalogram performed six weeks before her death.

For 30 years, since Herpes simplex was first isolated from the brain of a dying child (Smith, Lennette, and Reames, 1941), the virus has been known for its sporadic but virulent neurotropism. Occasionally giving rise to a fulminating infection of the newborn (Zuelzer and Stulberg, 1952; Nahmias, Dowdle, Josey, Naib, Painter, and Luce, 1969) and to a more benign meningoencephalitis at all ages (Afzelius-Alm, 1951; Adair, Gould, and Smadel, 1953; Meyer, Johnson, Crawford, Dascomb, and Rogers, 1960; Ross and Stevenson, 1961; Leider, Magoffin, Lennette, and Leonards, 1965; Olson, Buescher, Artenstein, and Parkman, 1967), it is also the usual cause of acute necrotizing encephalitis (Haymaker, 1949; van Bogaert, Radermecker, and Devos, 1955; Krücke, 1957; Jellinger, Poetsch, and Seitenberger, 1964), which may mimic cerebral tumour, abscess, subdural empyema, or tuberculous meningitis (Adams and Jennett, 1967). A few cases have been attributed to other infections, such as lymphocytic choriomeningitis (Scheid, Jochheim, and Stammier, 1955) and psittacosis (Duckett, Kelly, and Grant, 1963), or to an enterovirus (Heathfield, Filsworth, Wall, and Corsellis, 1967).

The natural history of necrotizing encephalitis due to Herpesvirus hominis (Haymaker, Smith, van Bogaert, and de Chenar, 1958; Brihaye, 1959; Carmon, Behar, and Beller, 1965; Bergouignan, Julien, Vital, and Duver, 1968; Jacoby, Blennerhasset, and Richardson, 1971) is divisible into stages. An influenza-like prodrome, when the mucocutaneous eruption may appear (Hunt and Comer, 1955), is shortly followed by symptoms of cerebral invasion—high fever, stupor, meningeal irritation, epilepsy, and hemiplegia. Sometimes the onset is indolent (Drachman and Adams, 1962; Blackwood, Dudgeon, Newns, and Phillips, 1966). Signs of an expanding mass (Wolf and Cowen, 1950; Bennett, ZuRhein, and Roberts, 1962; Pierce, Portnay, Leeds, Morrison, and Wehrle, 1964), commonly
frontal or temporal, often lead to surgical intervention (Adams and Jennett, 1967). Death occurs in the second or third week (Miller, Hesser, and Tompkins, 1966; Olson et al., 1967). Of those few who survive a month or more the majority face a prospect of permanent mental and physical handicap punctuated by frequent fits, as full recovery is exceptional (Meyer et al., 1960; Leider et al., 1965; Miller et al., 1966; Olson et al., 1967; Bergouignan et al., 1968; Miller and Ross, 1968).

The infection is diagnosed by brain biopsy (Dodge and Cure, 1956; Booth, Okazaki, and Gaulin, 1961; MacCallum, Potter, and Edwards, 1964) and confirmed by a fourfold or greater rise in serum antibody titre (Ross and Stevenson, 1961). A single titre > 512 is held to be diagnostic of active infection (Miller and Ross, 1968). Electron microscopy is a quick and reliable method of finding virus particles (Ryden, Moses, Ganote, and Beaver, 1965; Itabashi, Bass, and McCulloch, 1966; Delmas-Marsalet, Bergouignan, Castaing, du Pasquier, and Vital, 1968) but, being non-specific, must be supported by biological and serological tests (Harland, Adams, and McSeveney, 1967; Johnson, Olson, and Buescher, 1968).

Herpetic invasion of the brain—blood-borne, by local spread along olfactory and trigeminal nerves (Johnson, 1964), or from a latent source of infection (Leider et al., 1965; Rawls, Dyck, Klass, Green, and Herrmann, 1966)—is widespread but irregular. Areas of massive necrosis distort the frontal and temporal lobes, focal haemorrhages in the cortex often disrupting the subjacent white matter (Haymaker et al., 1958; Jellinger et al., 1964). Clusters of intranuclear (Cowdry, 1934) and cytoplasmic (Haymaker, 1949) inclusions establish the viral aetiology (Krücke, 1957). Most infections of older children and adults are with type I Herpesvirus hominis (Dowdle, Nahmias, Harwell, and Pauls, 1967) whereas type II virus is the usual cause of neonatal illness (Nahmias et al., 1969).


**CASE REPORTS**

Two cases are reported below of presumed necrotizing encephalitis due to infection with *Herpesvirus hominis*. In the first the virus was isolated by brain biopsy and the infection was confirmed by a diagnostic rise in the level of serum antibody. The second case also showed a significant rise in antibody titre but brain biopsy was unsuccessful. Serial scanning with technetium-99m pertechnetate was correlated with the clinical course of the illness, with angiographic and electroencephalographic findings, and with the effect of steroids and cytotoxic drugs.

**CASE 1**

A 6 year old boy was taken ill on 10 July 1970 with headache, fever, and sore throat. His family doctor prescribed ampicillin. During the evening of 12 July he had several convulsions lasting a quarter of an hour before he was admitted to a local hospital. His previous health had been excellent.

On admission he was stuporous with temperature 40°C, pulse 160 per minute, and blood pressure 120/80 mm Hg. No rash, ulcers, nor petechiae were seen. Heart, lungs, abdomen, and urine were normal. Slight neck stiffness was found. Pupils and fundi were normal, but the tone of all the limbs was increased,
tendon reflexes were absent, and both plantar responses were flexor.

A clear fluid was obtained at lumbar puncture, with 115 WBC (92% lymphocytes) and 12 RBC/c.mm, 42 mg protein, and 90 mg glucose/100 ml. Bacterial culture of the spinal fluid and of blood, throat swab, and urine, and virus culture of faeces were all negative. Cultures of spinal fluid for acid-fast bacteria were also negative. Other investigations included a radiograph of the chest which was normal and a blood count with haemoglobin 13.5 g/100 ml. WBC 14,000/c.mm (91% polymorphs, 9% lymphocytes), and sedimentation rate 10 mm in 1 hr. Serum electrolytes were normal, blood sugar 90 mg, and urea 20 mg/100 ml. Mantoux test was negative at 1:1,000.

Hydrocortisone was given intravenously, together with ampicillin, chloramphenicol, and sulphadiazine in doses adjusted for body weight (25 kg). Adrenocorticotrophin was begun on 17 July.

He remained unresponsive with an irregular fever swinging to 39-5° C until the evening of 17 July when he had two generalized convulsions in quick succession. A second lumbar puncture on the previous day had again shown a clear fluid with no increase of WBC but containing 250 RBC/c.mm, 35 mg protein, and 80 mg glucose/100 ml. An electroencephalogram (EEG) on the morning of 17 July showed generalized slow activity with a periodic delta focus in the left posterior temporal region (Fig. 1).

On 18 July he was transferred to Cardiff Royal Infirmary. He was comatose with neck rigidity, trismus, and positive Kernig sign. Persistent twitching of the right hand (epilepsia partialis continua) was seen. Pupils were dilated but reacted to light. The fundi were normal and corneal reflexes present. Rigidity of all four limbs was more marked on the right. Tendon reflexes were absent, the plantar responses both extensor. The skin and mucous membranes were normal. The liver edge was just palpable but no enlargement of spleen nor of lymph nodes was found. Intravenous diazepam controlled the focal epilepsy.

Left carotid angiogram (Fig. 2) on 20 July showed elevation of the middle cerebral vessels with stretching of the anterior temporal branch. The midline vessels were displaced to the right (Fig. 3). Cerebral scan on 21 July, one hour after intravenous injection of 2.5 mc technetium-99m pertechnetate, showed an area of increased uptake in the left posterior temporal region (Fig. 4) extending into the parietal area. On 22 July virus particles with the characteristics of *Herpesvirus hominis* were identified by electron microscopy in a biopsy specimen and the virus was later grown in tissue culture. By 28 July the complement fixation titre for *H. hominis* which had been <1:8 on 15 July had risen to 1:1024.

The therapeutic regime of steroids, adrenocorticotropic hormone and antibiotics was continued until 23 July. On that day cytosine arabinoside (cytarabine),

![Fig. 1](http://jnnp.bmj.com/content/35/5/569/fig-1)

*Fig. 1. Case 1. Electroencephalogram (17 July 1970). Generalized slow waves with periodic delta focus in left posterior temporal region.*
A. S. Bligh, C. M. Weaver, and C. E. C. Wells

FIG. 2. Case 1. Left carotid angiogram (20 July 1970). Lateral view showing elevation of middle cerebral vessels with stretching of anterior temporal branch.

FIG. 3. Case 1. Left carotid angiogram (20 July 1970). Anterior view showing displacement of midline vessels to right.

FIG. 4. Case 1. Isotope encephalogram (21 July 1970). Left lateral scan showing area of increased uptake in posterior temporal region.

FIG. 5. Case 1. Isotope encephalogram (4 August 1970). Right lateral scan showing multiple areas of abnormal uptake in frontal and parietal regions.

0·3 mg/kg/day (total 7·5 mg), was begun as an intravenous infusion, the dose being raised to 0·6 mg/kg/day (total 15 mg) from 24 July and to 1·8 mg/kg/day (total 45 mg) from 25 July. Penicillin and anticonvulsants were continued.

On 25 July he was lightly stuporous, sometimes opening his eyes and moving his limbs on command. He could now be fed by spoon. When his arms were raised passively they developed a coarse tremor and showed considerable rigidity. His improvement was maintained over the next 10 days and he became...
Isotope encephalography in the management of acute herpesvirus encephalitis

L Loatral 4.8 70.

FIG. 6. Case 1. Isotope encephalogram (4 August 1970). Anterior scan showing left-sided lesion which follows the curve of the vault. Faint area of abnormal uptake on the right.

FIG. 7. Case 1. Isotope encephalogram (4 August 1970). Left lateral scan showing migration of posterior temporal lesion to the periphery. Compare with Fig. 4.

Cytarabine was discontinued on 2 August but was restarted on 5 August when the third lumbar puncture showed 14 WBC and 41 RBC/c.mm, 88 mg protein, and 58 mg glucose/100 ml. The fluid was again clear and bacterial culture sterile.

The second brain scan on 4 August showed new areas of abnormal uptake in the right hemisphere (Fig. 5) and apparent migration of the left posterior

temporal lesion to the periphery (Figs 6 and 7). Left-sided twitching, partially suppressed by diazepam and phenytoin, began on 6 August, the EEG showing a right parietal ictal discharge (Fig. 8). At this stage he became less accessible, tremor increased and fever returned, and on 14 August he had several generalized fits. The EEG now showed flattening of the record in leads from the right hemisphere against a background of generalized slow activity. In a third scan on 21 August the left-sided lesion had increased in size and in intensity of uptake (Figs 9 and 10), although the EEG no longer showed a focal abnormality in this area.

Lumbar puncture on 22 August was normal. The fluid was clear with no increase of cells, 48 mg protein, and 65 mg glucose/100 ml. Intravenous idoxuridine was started on 24 August and five daily doses of 100 mg/kg (daily total 2.5 g) were given. No improvement was seen. He remained stuporous with frequent shuddering tremors, opening his eyes on painful stimulation only and continuously grimacing. Low remittent fever persisted. The optic discs remained flat, the right plantar response became flexor but the left was still extensor. Myoclonic jerks occurred in bursts which were independent of EEG changes. Reversal of sleep rhythm developed shortly before he returned to his local hospital on 3 September.

Throughout the period of cytarabine and idoxuridine therapy blood counts were performed regularly. During the first week of August he became anaemic and by 9 August the haemoglobin had fallen to 9.6 g/100 ml., with RBC 3.02 millions and WBC 3,100/c.mm (59% polymorphs, 39% lymphocytes, 1% monocytes, 1% eosinophils) and platelets 240,000 per c.mm. One unit of blood, group A Rh(D)+, was transfused on 12 August. Thereafter repeated counts were normal.

During the next three months consciousness gradually returned and the frequency of his epileptic fits diminished. By January 1971 he was able to sit unaided, to walk between bars, and to stand by supporting himself on the furniture. Ten months later he had not regained useful speech, could not feed himself, and had no proper control of sphincters. Major and minor epileptic fits continued to occur several times a day.

**CASE 2**

A 9 year old boy had been listless for a fortnight when he started to complain of headache and abdominal pain. His general practitioner found that he was febrile and prescribed paracetamol. Within 48 hours he had become drowsy and ampicillin was given. Three days later, on 11 October 1971, he was admitted to hospital.

On transfer to Cardiff Royal Infirmary on the evening of the same day he was conscious but inattentive, lay curled on his side and resented examination. His temperature was 38°C and pulse rate 82 per minute. Neck stiffness and Kernig's sign were found. He had a mild left hemiparesis, with sluggish reflexes and flexor plantar responses. Soon after his admission he had three small fits which were followed by deepening stupor and dense left hemiplegia with left extensor plantar response.

The spinal fluid was clear with resting pressure 180 mm. It contained 266 RBC and 154 WBC (mostly lymphocytes)/c.mm, 65 mg protein, and 70 mg glucose/100 ml. Bacterial culture was sterile and no organisms were seen on the Ziehl-Neelsen film. The peripheral blood showed a leucocytosis of 15,600/c.mm. The Mantoux test was negative at 1:1,000.

In the first EEG on 11 October diffuse slow activity at delta and sub-delta frequencies was seen, with greater amplitude in leads from the right hemisphere. There was no focus. Further records on 15 October and 20 October were similar. The first brain scan with technetium-99m pertechnetate on 12 October indicated a right occipitotemporal lesion which had become more pronounced in the second scan (Figs 11 and 12) on 15 October. Tissue from a right frontal brain biopsy on 12 October was unsuitable for laboratory examination and was not processed. Complement fixation tests for *Herpes simplex* showed a rise in titre from 1:16 on 11 October to 1:512 on 22 October. Culture of faeces for enteroviruses was negative.

Viral encephalitis was diagnosed on the clinical, EEG, and laboratory findings and, in view of the abnormal scan, herpes simplex was suspected. The red cell pleocytosis of the spinal fluid, which was a feature of the first case, was thought to be additional evidence in favour of this diagnosis (Miller et al., 1966). Although proof of herpesvirus infection was not forthcoming, the rising titre of serum antibody was highly significant (Ross and Stevenson, 1961; Miller and Ross, 1968).

Cytosine arabinoside (cytarabine) was given daily as an intravenous infusion from 12 to 18 October in a dose of 3-5 mg/kg/day. The total dose was 700 mg (weight 28-25 kg). In addition dexamethasone, 8 mg daily in divided doses, was given from 12 to 21 October, was then reduced and stopped on 23 October. Phenytoin was given throughout his stay in hospital.

By the evening of 13 October, 24 hours after chemotherapy and steroids had begun, he was conscious and afebrile but the left hemiplegia persisted. On regaining consciousness he was found to have a left homonymous hemianopia. He spoke to his mother on 16 October and took a few steps with sup-
Isotope encephalography in the management of acute herpesvirus encephalitis

FIG. 9. Case 1. Isotope encephalogram (21 August 1970). Posterior scan showing dense area of abnormal uptake lying just below the vault and predominantly left-sided.


FIG. 11. Case 2. Isotope encephalogram (15 October 1971). Right lateral scan showing occipitotemporal lesion.

FIG. 12. Case 2. Isotope encephalogram (15 October 1971). Posterior scan showing increased uptake of right occipitotemporal lesion.
FIG. 13. Case 2. Isotope encephalogram (2 November 1971). Right lateral scan showing diminished uptake of occiptotemporal lesion. Compare with Fig. 11.

port. Further improvement was delayed by sensory ataxia of his left side, affecting chiefly the hand and fingers. By 23 October he was able to identify objects in his left visual field but inattention hemianopia persisted. He was well enough to return home on 25 October. A third brain scan (Fig. 13) on 2 November showed that the right occiptotemporal lesion was still present, although the intensity of uptake was less. In the posterior scan (Fig. 14) the abnormal area lay superficially. A month later he was well and without abnormal physical signs. The scan on 1 December was normal.

In a previous case we had found that brain

FIG. 14. Case 2. Isotope encephalogram (2 November 1971). Posterior scan showing improvement of right occiptotemporal lesion. Compare with Fig. 12.

FIG. 15. Case 3. Isotope encephalogram (21 October 1968). Right lateral scan showing increased frontotemporal uptake.

scanning could provide a vital diagnostic clue when a patient suspected of carcinomatous, or of spongiform, encephalopathy was found at necropsy to have necrotizing encephalitis.

CASE 3
A 63 year old woman had a three-month story of progressive loss of memory followed by unsteadiness, tremor, and drowsiness. She had lost much weight. On admission to hospital she was afebrile but later ran an irregular fever up to 38.5°C, coinciding with clinical and radiological signs of pulmonary infection. General examination including the pelvis was otherwise normal.

Within days she became mute, earlier appearing confused and disoriented with almost total loss of recent memory and of retention and recall, and showing marked perseveration of both speech and action. She sat for long periods gazing vacantly about her, her attitude of repose frequently interrupted by myoclonic jerks. She was too apraxic to stand, even with help, and the static tremor of her arms became turbulent on movement. Pupils and fundi were normal, the plantar responses flexor. Bilateral grasp and pouting reflexes were present.

Lumbar puncture showed a mild pleocytosis of 25 WBC/c.mm but on repetition the count was normal. Although the protein content was not raised, the Lange curve was 'paretic'. The EEG which was recorded twice was marred by artefact and showed a continuous background of low-voltage slow activity. An alpha rhythm at 9 Hz was distinguishable in parts of the first record. Paired sera for viral antibodies, reported shortly before her death, showed an initial titre to herpes simplex of 1/128 falling after 24 days to 1/64. Areas of increased uptake over the frontal lobes (Figs 15, 16, 17) were reported in the brain scan. Further investigation was abandoned because of her rapid deterioration.

At necropsy she was found to have a terminal bronchopneumonia and her right kidney had been destroyed by tuberculosis which was considered quiescent. No carcinoma was found. The brain showed changes of an acute encephalitis with necrosis and haemorrhage extending throughout the frontal and temporal lobes. The occipital lobes were less severely affected and the cerebellum and brainstem appeared macroscopically normal. Histological sections confirmed the gross appearance of necrotizing encephalitis but no inclusion bodies were found and culture of brain tissue obtained at necropsy was negative for viruses.

DISCUSSION

THE PATHOLOGICAL LESION A necrotizing encephalitis is the typical post-mortem finding in patients dying of herpesvirus infection of the brain (Haymaker, 1949; van Bogaert et al., 1955; Krücke, 1957; Haymaker et al., 1958; Brihaye, 1959; Jellinger et al., 1964; Adams and Jennett, 1967). Although the virus enters and destroys nerve-cells, astrocytes, and oligodendroglia (Wolf and Cowen, 1950; Ryden et al., 1965; Chou and Cherry, 1967; Bergouignan et al., 1968) its part in causing necrosis and haemorrhage is less clearly defined. Krücke (1957) recognized oedema, haemorrhage, and necrosis as separate stages of an evolving process. A viral vasculitis (Carmon et al., 1965; Hughes, 1969) is a further factor but it is not constantly present (Adams and Jennett, 1967).

Recovery from necrotizing encephalitis can occur (Dodge and Cure, 1956; Leider et al., 1965; Olson et al., 1967; Miller and Ross, 1968) but is much less likely than in cases of milder illness. Whereas the mortality rate for all cases of
herpes encephalitis is under 50%, nearly three-quarters of those with necrotizing lesions die (Miller and Ross, 1968). In most recent reports of recovery an antiviral drug, idoxuridine, often combined with surgical decompression, has been given (Breeden et al., 1966; Marshall, 1967; Page et al., 1967; Bellanti, Guin, Grassi, and Olson, 1968; Nolan et al., 1970; Meyer et al., 1971; Rappel et al., 1971). Some evidence exists that milder cases of meningoencephalitis and of aseptic meningitis, which recover spontaneously, do not have necrotizing lesions (Miller and Ross, 1968), although the opportunity for pathological study is rare. They are clearly distinct from another group of cases, also without necrotizing lesions at necropsy, which have died of a fulminating infection before haemorrhage and necrosis have developed (Krücke, 1957; Bellanti et al., 1968).

**POSITIVE AND NEGATIVE SCANS** Positive brain scans occur only in cases of severe encephalitis and are not found in the majority of non-bacterial infections of the central nervous system (Davis and Tavera, 1966; Pedersen and Haase, 1970). Balfour et al. (1967) found that the abnormal pattern of uptake shown by their case of herpes encephalitis was unique in their experience of more than 4,000 brain scans. They contrasted a rounded zone, suggestive of an intratemporal mass, in the lateral scan with an elliptical area, curving under the vault, in the anterior and posterior scans. Their description resembles our own findings (Figs 1–14) and accords well with the illustrations of some isotope encephalograms which accompany recent case reports of herpes encephalitis (Halpern et al., 1970; Mishkin, 1970; Radcliffe et al., 1971).

Serial scanning of our first case showed that new lesions were emerging in the right hemisphere at a time of clinical remission (Figs 5 and 6) and anticipated by days focal epilepsy and other signs of relapsing infection. Such changes, however, should be viewed with caution as they may reflect no more than the physiopathology of repair, a similar progression being commonplace in the serial scans of infarction (Glasgow et al., 1965; Molinari et al., 1967; Usher and Quinn, 1969; Marshall and Popham, 1970). The increasing left occipitotemporal uptake of the second scan was wrongly interpreted as evidence of continuing infection and led to unnecessary therapy with idoxuridine. Similarly in our second case increasing density of uptake in the right occipitotemporal region (Figs 11 and 12) was seen after recovery had begun, subsequent scans showing resolution (Figs 13 and 15).

**CHEMOTHERAPY** The place of chemotherapy in herpes simplex infections of the nervous system was at first uncertain and its use was restricted to cases of suspected encephalitis from which herpesvirus had been isolated. Early enthusiasm for idoxuridine, chosen because of its success in treating herpetic keratitis (Leopold, 1965), was based on single case reports despite their authors’ caution over accepting such slender evidence (Breeden et al., 1966; Buckley and MacCallum, 1967; Evans, Gray, Miller, Jones, Weeks, and Wells, 1967; Marshall, 1967). Later experience, however, confirmed the anti-viral action of the drug which was effective across the blood-brain barrier. Both mortality and long-term morbidity were reduced if idoxuridine was given promptly at the onset of encephalitis (Nolan et al., 1970; Meyer et al., 1971; Rappel et al., 1971).

Cytosine arabinoside was introduced by Juel-Jensen (1970) for the treatment of generalized primary herpes and of severe herpes in patients with immunoparesis. One case of simian herpesvirus encephalitis, a rare but lethal disorder, also recovered (Juel-Jensen, 1970). Its use in herpes simplex (Herpesvirus hominis) encephalitis has not previously been reported. Other forms of therapy which have been advocated are surgical decompression (Adams and Jennett, 1967), an interferon inducer (Bellanti, Catalano, and Chambers, 1971), and steroids (Upton, Barwick, and Foster, 1971) despite the theoretical dangers which include suppression of interferon (Longson and Beswick, 1971). Cytarabine has the advantage of being instantly available whereas idoxuridine needs careful and laborious preparation.

Our first case was treated with cytarabine, using a small dose, and subsequently with idoxuridine. Steroids and adrenocorticotrophin had been given before his admission to our unit. The result was poor, the child remaining hemiplegic, epileptic, and grossly retarded 18 months after the conclusion of therapy. A similar result had been seen in the case previously reported.
from the unit (Evans et al., 1967) in which idoxuridine was started on the 56th day of illness. That patient is now resident in an institution for mentally retarded children, a severe epileptic with residual hemiplegia, and is liable to outbreaks of disturbed behaviour.

The second case was treated with cytarabine using a larger dose, similar to that for malignant disease, which was combined with dexamethasone as a means of reducing cerebral oedema. His response was prompt with a quick return to consciousness and early remission of hemiplegia and hemianopia. In the short follow-up period improvement has continued and all drugs, including anticonvulsants, have been withdrawn.

**Electroencephalography** In his monograph on encephalitis and encephalopathy Radermecker (1956) described a typical sequence of EEG changes in acute necrotizing encephalitis. The generalized slow activity of diffuse brain disease was followed by marked flattening of the record in some leads, particularly those from the temporal areas, and by the emergence of epileptic and periodic complexes. An unusual feature was the apparent improvement of the EEG at the time of clinical deterioration.

Upton and Gumpert (1970), writing on the EEG diagnosis of herpes simplex encephalitis, found that periodic complexes, often focal over the temporal region, appeared during the first fortnight of the illness thereafter subsiding without parallel clinical improvement. The EEG recorded on the ninth day of illness and illustrated in their paper (Upton and Gumpert, 1970) is strikingly similar to that on the eighth day of our case 1 (Fig. 1)—a pattern of focal complexes arising from the left temporal region. Still present on the 12th day the periodic focal discharge had disappeared by the 18th day, although the brain scan on the 28th day (Figs 9 and 10) showed a persisting lesion of this area.

In cases of cerebral infarction delay in obtaining a positive scan has been attributed to oedema which obstructs the vessels at the margin of the lesion (Usher and Quinn, 1969). When the oedema subsides, after 10 to 14 days, the vessels dilate, capillaries proliferate, and abnormal quantities of isotope are concentrated in and around the infarct. In necrotizing encephalitis decay of electrical activity over an area of increased vascularity, as shown by the scan, may well indicate a region of almost total neuronal death. This may be the explanation for the series of events which were recorded clinically, electrically, and radiologically in the left posterior temporal region of our first case.

Dr. A. D. Evans provided the reports on virus serology and on virus isolation and culture, and Dr. B. H. Knight the account of the necropy findings on case 3.

**REFERENCES**


Isotope encephalography in the management of acute herpesvirus encephalitis