Use of cerebrospinal fluid drawn at pneumoencephalography in the study of monoamine metabolism in man

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SYNOPSIS Concentrations of homovanillic acid (HVA) and 5-hydroxyindoleacetic acid (5-HIAA) were significantly higher in CSF obtained after injection of air during pneumoencephalography (PEG) than in lumbar CSF, as drawn before the injection. There was a high correlation between levels in the 'mixed' and lumbar samples of CSF in the case of each of the two acids. The concentration of lumbar HVA, but not that of 5-HIAA, was negatively correlated with CSF pressure. 5-HIAA levels were low in both samples of CSF in a group of epileptics, by comparison with controls. In two patients with Kufs disease and in one with Niemann-Pick disease, the concentration of HVA was very low in the lumbar sample. The application of a standardized PEG technique in the study of monoamine metabolism in man is suggested.

After the demonstration of 5-hydroxyindoleacetic acid (5-HIAA) (Ashcroft and Sharman, 1960) and homovanillic acid (HVA) (Andén et al., 1963) in human cerebrospinal fluid (CSF), an increasing number of reports on the CSF levels of these acidic metabolites of monoamines has been appearing in the literature. In some of these studies ventricular fluid obtained during diagnostic or therapeutic surgery has been used. This work has provided useful information on the cerebral roles of the two amines as detected by significant variation from control levels of the concentrations of their respective metabolites in several diseases, mainly neurological (Ashcroft and Sharman, 1960; Guldberg et al., 1967; Anderson and Roos, 1969; Papeschi et al., 1972). Information on the effects of drugs has also been obtained (Papeschi et al., 1972).

In most studies, however, particularly those concerned with psychiatric patients, investiga-

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information on the concentration of these metabolites in a variety of neuropsychiatric diseases would be obtained.

METHODS

Thirty-five patients undergoing diagnostic PEG at the Montreal Neurological Hospital were included in this study, without previous selection. During this work, lumbar CSF from two cases with Kufs disease became available and the results are reported here. Patients with evidence of obstruction of CSF flow were excluded from this series; those results have been reported separately (Garelis and Sourkes, 1973).

After the evening meal patients did not have any food until the procedure had been completed on the following day (generally before noon). One-half to one hour before PEG was started preparatory medication was given. In most cases this consisted of codeine 60 mg, dimenhydrinate (Gravol) 50 mg, and diazepam (Valium) 10 mg, intramuscularly. Occasionally, sodium pentobarbitone (100 mg) was substituted for diazepam.

After the lumbar puncture had been performed, CSF pressure was recorded in the sitting position. For routine diagnostic purposes 3–4 ml of fluid were kept, then the first sample (‘lumbar’) for this study was collected. At this time, air was fractionally injected through the puncture needle, and CSF was collected through the needle. At the end of the procedure, when all the air had been injected, the second (‘mixed’) specimen was taken. Samples were rapidly frozen and kept at −20° C until analysed.

HVA and 5-HIAA were measured fluorimetrically in the same sample, as described by Papeschi and

### TABLE 1

<table>
<thead>
<tr>
<th>Diagnosis</th>
<th>Lumbar HVA</th>
<th>Mixed HVA</th>
<th>Lumbar 5-HIAA</th>
<th>Mixed 5-HIAA</th>
</tr>
</thead>
<tbody>
<tr>
<td>Controls</td>
<td>43.1 ± 3.5 (10)</td>
<td>65.9 ± 6 (11)*</td>
<td>32.1 ± 2.4 (16)</td>
<td>44.9 ± 4.8 (16)†</td>
</tr>
<tr>
<td>Epilepsy</td>
<td>42.8 ± 4.5 (12)</td>
<td>60.5 ± 6.9 (12)</td>
<td>24.6 ± 2 (12)‡</td>
<td>32.9 ± 5.1 (12)</td>
</tr>
<tr>
<td>Brain-stem encephalopathy</td>
<td>32.25 ± 8.4 (4)</td>
<td>37.5 ± 8.7 (4)‡</td>
<td>34.7 ± 1.9 (4)</td>
<td>34.3 ± 3.11 (4)</td>
</tr>
<tr>
<td>Kufs disease</td>
<td>Undetectable</td>
<td>—</td>
<td>35</td>
<td>—</td>
</tr>
<tr>
<td>Niemann-Pick variant</td>
<td>13</td>
<td>76</td>
<td>32</td>
<td>—</td>
</tr>
<tr>
<td>Possible Parkinsonism</td>
<td>36</td>
<td>62</td>
<td>27</td>
<td>41</td>
</tr>
<tr>
<td>Schizophrenia</td>
<td>—</td>
<td>153</td>
<td>15</td>
<td>90</td>
</tr>
</tbody>
</table>

* Different from lumbar at P < 0.005.
† Different from lumbar at P < 0.025.
‡ Different from controls at P < 0.025.

### TABLE 2

<table>
<thead>
<tr>
<th>Age, sex (yr)</th>
<th>Type of seizure</th>
<th>Drug* (mg/day)</th>
<th>HVA Lumbar</th>
<th>HVA Mixed</th>
<th>5-HIAA Lumbar</th>
<th>5-HIAA Mixed</th>
</tr>
</thead>
<tbody>
<tr>
<td>18 F</td>
<td>Temporal</td>
<td>Phenytoin 200, phenobarbitone 60</td>
<td>69</td>
<td>111</td>
<td>25</td>
<td>38</td>
</tr>
<tr>
<td>22 M</td>
<td>Temporal</td>
<td>Primidone 200, ethosuximide 750</td>
<td>44</td>
<td>44</td>
<td>23</td>
<td>8</td>
</tr>
<tr>
<td>22 M</td>
<td>Temporal</td>
<td>Methylphenobarbitone 300</td>
<td>41</td>
<td>45</td>
<td>34</td>
<td>36</td>
</tr>
<tr>
<td>21 M</td>
<td>Temporal</td>
<td>Phenytoine 120, chloral 500</td>
<td>26</td>
<td>50</td>
<td>32</td>
<td>36</td>
</tr>
<tr>
<td>52 M</td>
<td>Temporal</td>
<td>Phenytoine 120</td>
<td>50</td>
<td>82</td>
<td>16</td>
<td>36</td>
</tr>
<tr>
<td>39 M</td>
<td>Temporal</td>
<td>—</td>
<td>36</td>
<td>41</td>
<td>18</td>
<td>21</td>
</tr>
<tr>
<td>62 M</td>
<td>Temporal</td>
<td>—</td>
<td>50</td>
<td>56</td>
<td>29</td>
<td>45</td>
</tr>
<tr>
<td>15 M</td>
<td>Corticocerebral</td>
<td>Primidone 750</td>
<td>31</td>
<td>51</td>
<td>33</td>
<td>82</td>
</tr>
<tr>
<td>18 M</td>
<td>Temporal—grand mal (withdrawal?)</td>
<td>Phenytoin 300, phenobarbitone 90</td>
<td>43</td>
<td>88</td>
<td>13</td>
<td>34</td>
</tr>
<tr>
<td>50 F</td>
<td>Grand mal</td>
<td>Phenytoin 300, phenobarbitone 30, primidone 750</td>
<td>72</td>
<td>81</td>
<td>23</td>
<td>28</td>
</tr>
<tr>
<td>39 M</td>
<td>Grand mal</td>
<td>Phenytoine 30, ethosuximide 1000</td>
<td>27</td>
<td>44</td>
<td>28</td>
<td>33</td>
</tr>
<tr>
<td>26 F</td>
<td>Grand mal</td>
<td>Phenytoin 240</td>
<td>25</td>
<td>33</td>
<td>21</td>
<td>22</td>
</tr>
</tbody>
</table>
McClure (1971). Clinical information and PEG findings were always obtained after the biochemical determinations had been performed.

For statistical analysis, subjects were grouped according to diagnostic category. Criteria for the selection of controls were: (1) absence of any disease where changes in serotonin or dopamine metabolism have been implicated, and (2) non-use of medication known to affect monoamine metabolism—for example, sedatives, neuroleptics. The diagnoses in the control group included the following: headache, facial pain, hysterical neurosis, spastic hemiparesis, optic atrophy, arterial aneurysm, chromophobe adenoma.

RESULTS

HVA and 5-HIAA concentrations in the lumbar and mixed samples are shown in Table 1 along with clinical diagnoses.

Acid levels of the control group were significantly higher in the mixed than in the lumbar specimen. This was also true for all other subjects, with the exception of cases of brain-stem encephalopathy (BSE).

The epileptic patients had HVA concentrations in their CSF samples similar to control values. On the other hand, 5-HIAA was lower in both lumbar and mixed fluid than in the corresponding control samples; this difference reached statistical significance in the lumbar CSF (Table 1). The data for the epileptic patients, including clinical information, are set out in Table 2.

No HVA could be detected in the lumbar CSF of the two siblings with Kufs disease. HVA was also low in the first sample of the patient with Niemann-Pick disease, but normal in the mixed one. One patient possibly suffering from Parkinsonism had normal concentrations of both acids. A schizophrenic subject had very high HVA and 5HIAA in the mixed CSF, but low 5-HIAA in the lumbar sample.

Correlation coefficients between pairs of the investigated variables were calculated and are shown in Table 3. Concentrations of the acids were highly correlated in both the lumbar and the mixed specimens in the control group, but not in the patients with epilepsy or BSE. The correlation between the metabolite concentrations in the lumbar CSF of our controls was in close agreement with data obtained in healthy volunteers (Gottfries et al., 1971). There was a significant negative correlation between CSF pressure and lumbar HVA but not 5-HIAA.

Because of the gradient of concentrations of metabolites, higher values being found in the upper CSF compartments (Table 1; Garelis and Sourkes, 1973), the amount of injected air or the total volume of withdrawn CSF might yield a positive correlation with the concentration of the metabolites in the mixed sample. However, the present data provided no basis for such correlations (Table 3).

**DISCUSSION**

It has been suggested that the injection of air during PEG results in concentrations of metabolites that would reflect ventricular rather than lumbar levels (Moir et al., 1970). This would be due to mixing of CSF of various compartments by the ascending air, and to disturbance of the normal gradient from the lateral ventricle through the cisterna magna to the lumbar level. Few investigations have taken advantage of PEG to study physiological or clinical relationships of monoamine metabolites. There is only one study directly comparing concentrations of 5-HIAA before and after the injection of air in

**TABLE 3**

**CORRELATIONS BETWEEN MEASURED VARIABLES**

<table>
<thead>
<tr>
<th>Variables</th>
<th>Subjects</th>
<th>N</th>
<th>R</th>
<th>P*</th>
</tr>
</thead>
<tbody>
<tr>
<td>HVA vs 5-HIAA, in lumbar CSF</td>
<td>Controls</td>
<td>10</td>
<td>0.771</td>
<td>&lt;0.005</td>
</tr>
<tr>
<td></td>
<td>Epilepsy</td>
<td>12</td>
<td>0.238</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>BSE†</td>
<td>4</td>
<td>0.840</td>
<td>NS</td>
</tr>
<tr>
<td>HVA vs 5-HIAA, in mixed CSF</td>
<td>Controls</td>
<td>11</td>
<td>0.767</td>
<td>&lt;0.005</td>
</tr>
<tr>
<td></td>
<td>Epilepsy</td>
<td>12</td>
<td>0.126</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>BSE</td>
<td>4</td>
<td>0.679</td>
<td>NS</td>
</tr>
<tr>
<td>Lumbar HVA vs CSF pressure</td>
<td>All cases†</td>
<td>26</td>
<td>-0.474</td>
<td>&lt;0.02</td>
</tr>
<tr>
<td>Lumbar 5-HIAA vs CSF pressure</td>
<td>All cases</td>
<td>32</td>
<td>0.21</td>
<td>NS</td>
</tr>
<tr>
<td>HVA in mixed CSF vs total volume of injected air (range = 40-125 ml)</td>
<td>Controls</td>
<td>16</td>
<td>0.223</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>Epilepsy</td>
<td>12</td>
<td>0.086</td>
<td>NS</td>
</tr>
<tr>
<td>5-HIAA in mixed CSF vs total volume of injected air (range = 40-125 ml)</td>
<td>Controls</td>
<td>16</td>
<td>0.213</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>Epilepsy</td>
<td>12</td>
<td>0.134</td>
<td>NS</td>
</tr>
<tr>
<td>HVA in mixed CSF vs total volume of CSF withdrawn (range = 25-65 ml)</td>
<td>Controls</td>
<td>11</td>
<td>0.363</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>Epilepsy</td>
<td>12</td>
<td>0.419</td>
<td>NS</td>
</tr>
<tr>
<td>5-HIAA in mixed CSF vs total volume of CSF (range = 25-65 ml)</td>
<td>Controls</td>
<td>16</td>
<td>0.216</td>
<td>NS</td>
</tr>
<tr>
<td></td>
<td>Epilepsy</td>
<td>12</td>
<td>0.600</td>
<td>&lt;0.05</td>
</tr>
</tbody>
</table>

* Probabilities. NS represents P > 0.05.
† BSE = Brain-stem encephalopathy.
†† All cases are pooled where compared variables did not differ significantly.
three patients; the concentration of 5-HIAA in the mixed sample was approximately double the lumbar level (Eccleston et al., 1970). In two more reports from the same group, similar values for 5-HIAA were found in the mixed sample, but lumbar fluid was not taken (Ashcroft and Sharman, 1960; Ashcroft et al., 1966).

In our control group, concentrations of 5-HIAA in both samples were higher than those reported by the Edinburgh investigators. The mixed/lumbar ratio for 5-HIAA however, was substantially smaller in our case. Several methodological differences might account for this discrepancy. The amount of air injected and the volume of CSF withdrawn were much greater in the present study. The injection of large amounts of air, much of which was directed to subarachnoid space outside the ventricles, coupled with the withdrawal of a large volume of CSF and the ensuing drastic changes in hydrostatic pressure would result in a downward current of fluid not only from the ventricles, but also from the subarachnoid space surrounding the brain. Concentrations of the metabolites in the latter space were not known, but might be relatively low, inasmuch as transport of the acids from the central nervous system is located mainly in the choroid plexuses (Ashcroft et al., 1968; Cserr and Van Dyke, 1971; Forn, 1972). Thus, the mixing process affects compartments with different mean concentrations of the metabolites, and is itself affected by the volume of air injected and fluid displaced. The lack of significant correlation between these two variables on the one hand, and acid concentrations in the mixed samples on the other (Table 3), was not surprising.

Another factor which could explain our failure to obtain the high concentrations of metabolites characteristic of the ventricular and cisternal compartments in the mixed samples was the effect of premedication. The patients in the study of Ashcroft et al. (1966) received only paracetamol orally, whereas all our patients had had heavy preparatory medication (see Methods section). Antihistamines (Coyle and Snyder, 1969), barbiturates (Corrodi et al., 1966, 1967), and benzodiazepines (Corrodi et al., 1967; Taylor and Laverty, 1969; Chase et al., 1970; Wise et al., 1972; Papeschi et al., 1972) are known to influence monoamine metabolism. Pharmacologically induced changes in serotonin or dopamine metabolism are evident in the cisternal 5-HIAA (Andersson and Roos, 1968; Eccleston et al., 1968) and HVA (Pletscher et al., 1967) in about one hour, but much later in the lumbar fluid. The interval between injection of premedication and collection of the mixed sample in the present study was 1--1.5 hours and less than one hour for the lumbar specimen. Under these conditions, therefore, premedication would affect the mixed, but not the lumbar sample.

Moreover, a possible effect of these drugs on the transport mechanisms and local hydrostatic pressure cannot be excluded. It would be difficult to predict, from the available data, what the net effect of the drug combinations used in this study on the acid levels in CSF would be.

The somewhat sharper gradient for HVA than 5-HIAA on going from the mixed sample to the lumbar CSF (Table 1) could be accounted for by the fact that 5-HIAA, but not HVA, in the CSF originates to a substantial extent from the spinal cord (Garelis and Sourkes, 1973; Young et al., 1973); consequently, mixing during PEG would disturb the 5-HIAA gradient less than that of HVA.

Low ventricular (Papeschi et al., 1972) and lumbar concentrations of HVA (Bernheimer et al., 1966; Barolin and Hornykiewicz, 1967) have been reported in some epileptic patients. This was not the case in our series in either the mixed or the lumbar sample. It is difficult to account for the apparent discrepancy in regard to HVA. However, it should be recalled that half the patients in the series of Barolin and Hornykiewicz (1967), mainly tumour epilepsies, had normal values.

In contrast with the result with HVA, low 5-HIAA levels were found in both specimens; the difference from controls reached significance for the lumbar fluid (Table 1). In the recent study from this laboratory (Papeschi et al., 1972), three cases of temporal lobe epilepsy had low ventricular 5-HIAA. This finding could not be further evaluated at the time for lack of adequate controls, but the concentrations were much lower than reported ventricular, or even normal cisternal, levels (Moir et al., 1970; Garelis and Sourkes, 1973). Moreover, preliminary results obtained by Shaywitz et al. (1973) showed a
decreased accumulation of 5-HIAA after pro-
benecid in epilepsy.

Epilepsy is not a homogeneous disease. Grouping of all patients with convulsions, regardless of aetiology, localization of the lesion, type and frequency of seizures, or concomitant psychiatric symptoms, may obscure possible associations of the biogenic amines with convul-
sive disorders. More important, the effect of antiepileptic medication has scarcely been studied. Barolin and Hornykiewicz (1967) pro-
vided evidence that low lumbar HVA in some of their patients was not due to the effect of diphenylhydantoin. However, the effect of chronically administered combinations of anti-
convulsants on CSF levels of monoamine metabolites is currently unknown. It can be seen from Table 2 that variations in the parameters mentioned above make practically each patient in our group different from the others.

We have no ready explanation for the absence of an increase of concentrations of the acids in the mixed fluid of the patients with BSE. Since most of the serotoninergic neurones are in the brain-stem it might be postulated that 5-HIAA would be low in the mixed sample because of destruction of these neurones. In the lumbar sample, this decrease could be compensated by a contribution from spinal interneurones (Post et al., 1973). This mechanism, however, should leave HVA unaffected. Some of these patients were probably suffering from multiple sclerosis, but this diagnosis was avoided in the absence of clear-cut recurrent episodes. Low concentrations of HVA and/or 5-HIAA in multiple sclerosis have recently been reported (Johansson et al., 1972).

The virtual absence of HVA in the two siblings with Kuf's disease was an unexpected finding. Although these patients are known to manifest occasionally extrapyramidal symptoms (Sourkes, 1962), decreased dopamine metabolism has never, to the best of our knowledge, been implicated in any of the lipidoses. Our two patients did not have extrapyramidal symptoms.

The patient suffering from a variant of Niemann-Pick disease also had low HVA in the lumbar sample. But the high level in the mixed fluid suggested that reduced flow of the acid towards the lumbar space, rather than a de-
fiency in dopamine metabolism was responsible for the low lumbar levels. Although the present data are few in number, the results indicate the need for further studies of dopamine metabolism in the lipidoses.

The possibility of Parkinson's disease in the next patient (Table 1) was raised because of the presence of an obscure tremor of recent onset. Notwithstanding the doubts about the diagnosis, it was not surprising that HVA was normal in this patient, for it has been shown (Papeschi et al., 1972) that low HVA levels are particularly correlated with akinesia; this was absent in the present patient.

Concentrations of both acids in the mixed fluid of the schizophrenic patient are the highest in the present series. Bowers (1969) has reported a case of schizoaffective psychosis with very high levels of both acids and provided evidence for a defective transport mechanism in that case. This possibility can be excluded in our patient because in such a condition, lumbar 5-HIAA would also be high. Low concentrations of 5-HIAA in schizophrenia have been reported (Ashcroft et al., 1966; Bowers et al., 1969) but other investigators were not able to confirm this finding (Persson and Roos, 1969; Rimón et al., 1971).

The finding that the hydrostatic pressure of the CSF is related to lumbar levels of HVA negatively may seem surprising, as it is known that high intracranial pressure (hydrocephalus) is associated with very high concentrations of both HVA and 5-HIAA. However, this is pre-
sumed to result from blockade of the transport mechanism (Andersson and Roos, 1968). In this study there were no cases with signs of hydro-
cephalus; the pressure range was within normal limits. Under these conditions, there is no question of an effect on the transport mechanism.

Local hydrostatic pressure is known to affect the flow of CSF (Davson, 1967). Most of or all the lumbar HVA comes from higher levels via the cisterna magna (Curzon et al., 1971; Garelis and Sourkes, 1973; Post et al., 1973). Changes in the cisterno-lumbar flow therefore, induced by local variations in pressure, would affect the lumbar levels of HVA. On the other hand, a sub-
stantial proportion of 5-HIAA originates in the spinal cord (Garelis and Sourkes, 1973). Conse-
quently, small variations in CSF flow from the cisterna magna would affect the total lumbar
concentrations to a far smaller extent. This could explain the absence of a significant effect of pressure on lumbar levels of 5-HIAA (Table 3).

One cannot exclude the possibility that higher pressures favour more rapid movement of water from the lumbar CSF compartment and bulk transport of solutes, like HVA, along with it.

On the question of the proportion of 5-HIAA originating in the spinal cord, our original estimate was based on controls selected only on the basis of absence of obstruction of CSF flow (Garelis and Sourkes, 1973). Using the much more refined present group of controls, 5-HIAA of spinal origin is estimated to account for 39% of the lumbar concentration. The recent paper by Weir et al. (1973) of a study in experimental animals reports a similar value.

The results of the present work show how one can take advantage of PEG as a very useful technique in the study of monoamine metabolism in man. In this study we deliberately avoided any attempt to standardize the conditions of obtaining samples because our aim was to study the effect of a range of different conditions. However, it is conceivable that injection of a small amount of air directed into the ventricles, followed by withdrawal of a small quantity of CSF under standardized conditions, could be effective in providing enough mixing to give a reliable indication of the concentrations of the acidic metabolites in the ventricular system, and thus to provide some reflection of amine metabolism in the brain. Under the above conditions premedication, clearly a disadvantage, would not be indispensable. Nor would the procedure be more distressing to the patients than the two lumbar punctures routinely employed in the probenecicd technique of assessing the metabolite concentrations.

The importance of seeking new techniques needs to be stressed, as means of obtaining decisive information about the state and metabolism of biogenic amines in the healthy and ailing human brain during life. Information of this kind ought to help greatly in clarifying the role of these substances in important diseases where they have already been implicated: in schizophrenia, manic-depressive psychosis, and diseases of the basal ganglia.

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