Short report

Lymphocyte subsets at different stages of subacute sclerosing panencephalitis: a study with monoclonal antibodies

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SUMMARY  Lymphocyte subsets in cerebrospinal fluid (CSF) and peripheral blood were studied using monoclonal antibodies, in patients with subacute sclerosing panencephalitis, eight of whom were at stage 2 and seven at stage 4. Eighteen subjects affected with non immunological diseases constituted the controls. Regardless of the stage, patients with subacute sclerosing panencephalitis had lower percentages of OKT3+ (pan-T) cells in both CSF and peripheral blood, with an increase of OKT8+ cells (B cells, macrophages and active T cells) in peripheral blood. A difference was found in the proportion of OKT4+ (helper-inducer) and OKT8+ (suppressor/cytotoxic) cells in relation to the stage, the most striking finding being a significant decrease of OKT8+ with an increase of T4/T8 ratio in peripheral blood at an early stage.

The persistence of virus-infected cells in the central nervous system (CNS), in spite of high levels of serum and cerebrospinal fluid (CSF) antibodies to measles antigens, suggests that the immune system might be involved in the pathogenesis of subacute sclerosing panencephalitis.1,2 We have studied lymphocyte subpopulations in CSF and peripheral blood from patients with subacute sclerosing panencephalitis.3,4 The percentages of all T lymphocytes (identified as E-rosette forming cells, E-RFC) and of T lymphocytes bearing IgG Fc receptors (identified as EA-rosette forming cells, EA-RFC, or TG cells, or "suppressors"), which represent the majority of lymphocytes in the CSF from subjects affected with "non immunological diseases" (NID), were found to be significantly lower in patients with subacute sclerosing panencephalitis.3,4 We recently investigated the antigenic phenotype of CSF lymphocytes using monoclonal antibodies.5 These studies showed that in non immunological diseases (a) 90% of CSF cells were T lymphocytes, as they reacted with OKT3, and (b) about 30% were T suppressor cells, identified by OKT8, that is a much lower percentage than that of TG cells.

We now report our findings regarding: (a) the evaluation of CSF lymphocyte subsets in patients with subacute sclerosing panencephalitis by monoclonal antibodies, and (b) the comparison of immunological parameters in patients at early and late stages of the disease.

Material and method

Patients  Twelve patients affected with subacute sclerosing panencephalitis (seven males and five females, mean age 9, range 4–12 years) were studied. Diagnosis was based on clinical, electroencephalographic and laboratory (serum and CSF antimeasles antibody titre, CSF IgG content and CSF protein electrophoresis) findings. CSF cell content was in the normal range in all cases (0·6–6·0/mm³). Eight patients were studied at clinical stage (according to Jabbour et al6), 14 days to 5 months (mean 1·8 months) after the first signs of the disease. Seven patients (three of whom had already been studied at stage 2) were studied at clinical stage 4, from 4 to 75 months (mean 23·5 months) after the onset of subacute sclerosing panencephalitis. All the latter had been treated for a period of 7–65 (mean 13·7) months with inosiplex. Thymostimuline was administered to three of them for a short period several months before the date of the immunological
examinations. No patient at stage 2 was treated. Eighteen subjects (nine males and nine females, mean age 27, range 4–52 years), affected with non immunological diseases, were studied as controls: eight were affected with epilepsy, four with spondylotic myelopathy, three with neurosis and three with hydrocephalus.

**CSF and peripheral blood lymphocytes** Cells were obtained according to the method previously described. Monoclonal antibodies. OKT3 (all-T), OKT4 (T helper-inducer), OKT8 (T suppressor-cytotoxic), OKIa (DR framework: B cells, macrophages and some activated T cells) were obtained from Ortho Diagnostico System (USA). The indirect fluorescence method was followed.

**Results**

Data on mononuclear cells identified in CSF and peripheral blood from subacute sclerosing panencephalitis and non immunological diseases patients are shown in the table. Significant findings in all the patients with subacute sclerosing panencephalitis are:

(a) a lower percentage of OKT3+ cells in both CSF and peripheral blood, of OKT4+ cells in CSF and of OKT8+ cells in peripheral blood; (b) an increased percentage of OKIa+ cells in peripheral blood. The comparison between patients at stage 2 and 4 shows:

(a) in CSF, a lower percentage of OKT8+ cells in stage 4; (b) in peripheral blood, a lower percentage of OKT8+ cells with increased T4/T8 ratio in stage 2 and reduced OKT4+ cells in stage 4.

**Discussion**

The evaluation of CSF and peripheral blood lymphocytes using monoclonal antibodies OKT3 in patients with subacute sclerosing panencephalitis, compared with patients with non immunological diseases, confirms the decrease in T cells previously observed using the E-rosette method. Results with EA-RFC, bearing FcIgG receptors and considered to have suppressor capability, differ slightly from the present finding with OKT8 (a marker of T suppressor cells), since EA-RFC were reduced in both peripheral blood and CSF in subacute sclerosing panencephalitis. Moreover, percentages of EA-RFC in CSF were always higher than those of OKT8+ cells, in both non immunological diseases and subacute sclerosing panencephalitis patients (it should be remarked that in CSF the percentage of B cells is very low, about 2%; therefore most cells bearing IgG Fc receptors are T; see ref 7). This is probably due to the fact that IgG-Fc receptors (EA rosette method) and OKT8 monoclonal antibodies both identify cells having suppressor capability, but neither marks all of them (reviewed in ref 8).

The behaviour of lymphocyte subsets at different stages of the disease deserves some consideration. The constant decrease of OKT3+ (pan-T) cells appears in some cases to be due mainly to reduced OKT4+ cells (OKT8+ being unchanged) and in others to reduced OKT8+ cells. In CSF, the decrease of OKT4+ at stage 2 (OKT8+ being unchanged) contrasts with the decrease of OKT8+ at stage 4, while the contrary occurs in peripheral blood. Findings at stage 4 are not likely to be due to inosiplex treatment, since the drug is not known to have similar effects on lymphocytes.

The reduction in CSF of some cell subsets, unchanged in peripheral blood, might be due either to reduced passage to the CSF or to sequestration and/or destruction in the CNS. While no data support the possibility of selective reduced passage to the CSF, a selective increase of some lymphocyte subpopulations in CNS is reported in multiple sclerosis and in experimental allergic encephalomyelitis. The behaviour of CSF lymphocytes in subacute sclerosing panencephalitis might reflect a similar condition: that is, the decrease of OKT4+ cells at stage 2 and of

<table>
<thead>
<tr>
<th>Table</th>
<th>Percentages of cells reacting with monoclonal antibodies from CSF and peripheral blood from patients with subacute sclerosing panencephalitis (SSPE) and from patients affected with non immunological diseases (NID)</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>SSPE stage 2</strong></td>
<td><strong>SSPE stage 4</strong></td>
</tr>
<tr>
<td>OK3</td>
<td>OK4</td>
</tr>
<tr>
<td>63.9 ± 14.2 (8)</td>
<td>43.2 ± 4.9 (4)</td>
</tr>
<tr>
<td>30.7 ± 11.5 (7)</td>
<td></td>
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<tr>
<td>1.9 ± 0.4 (4)</td>
<td></td>
</tr>
</tbody>
</table>

Percentages of cells reacting with monoclonal antibodies, mean ± standard deviation (number of cases in brackets).

Statistically significant differences (Student's t test: *p < 0.05, †p < 0.01, ‡p < 0.001); All SSPE vs NID: CSF T3*, T4*, PB T3†, T8†, Ia*. Stage 2 vs NID: CSF T3†, T4*, PB T8†, T4/T8*, Ia*. Stage 4 vs NID: CSF T3*, T8†, PB T3*, T4*. Stage 2 vs stage 4: CSF T8*, PB T8*, T4/T8*.

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OKT8+ at stage 4 might be the result of selective sequestration in the central nervous tissue. The findings in peripheral blood indicate a more generalised change of the cellular immune system in subacute sclerosing panencephalitis. T-cells appear to be on the whole reduced, while 1a + cells (which include B lymphocytes, macrophages and some activated T cells) increase proportionally. The striking decrease of OKT8+ cells and the consequent increase of the T4/T8 ratio at the early stage of subacute sclerosing panencephalitis may suggest some similarities in the behaviour of the immune response to the acute phases of multiple sclerosis, in which OKT8+ reduction is also reported.11 A decrease of total T cells in peripheral blood has also been found by others.12 13 Moreover, Vainiene et al13 found a progressive decrease in T cells with FcIgG receptors in patients with a rapid course of subacute sclerosing panencephalitis, while they reported a progressive increase in stable cases. This appears to agree with our results of a progressive increase of OKT8+ cells from the “evolutive” stage 2 to stage 4, a condition of disease stability.

A few other abnormalities were found in cases of subacute sclerosing panencephalitis,11 12 13 although they were not always correlated with the stage of the disease. The significance of such abnormalities in the immune system needs to be explained. It should be noted that measles virus is found to have a depressing effect on T cells.1 Therefore, it remains to be ascertained whether the decrease in T cells in subacute sclerosing panencephalitis might be simply an epiphenomenon, due to the persistence of such a viral effect, or if it is involved in the pathogenetic mechanism of the disease. Supporting this latter hypothesis is the occurrence of more acute measles encephalitides (measles inclusion body encephalitis),15 and of subacute sclerosing panencephalitis16 in subjects with marked depression of cellular immunity.

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References

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