34th Kinki Neuropathology Conference
(Prof. Scheithauer's slide session)

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Chairmen: Yasuaki Nakashima (Kyoto University Hospital, Surgical pathology)
Keiji Kawamoto (Kansai Medical University, Neurosurgery)

Special Lecture
"Newly Proposed WHO Histological Typing of the Central Nervous System"

Malignant lymphoma or multiple sclerosis?

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Radiological and neuropathological findings of a case with two possible but different diagnoses, malignant lymphoma (ML) or multiple sclerosis (MS), was presented. At first a 40 y.o housewife developed typical symptoms of intracranial tumor with raised intracranial pressure and space-occupying mass in the rt frontal lobe on CT and MRI. But surgical specimens (1991. 4. 25.) disclosed only inflammatory appearance, consisting of proliferating astroglia interspersed by numerous foamy macrophages, and perivascular lymphocyte cuffings. The feature was similar to biotical findings of multiple sclerosis mimicking primary brain tumor described by Hunter et al. in 1987. Prior to operation, she had received glucocorticoid therapy (12mg q. d. dexamethasone for 6 days), which led remarkable improvement of neurological state. So the clinical side insisted on the diagnosis of malignant lymphoma obscured by glucocorticoid therapy because of (1) the patient’s past history of ML in the axillary lymph node 9 years previously, (2) relative rarity of MS in Japan, and (3) glucocorticoid administration prior to surgery. Rapid regression

Prof. Bernd Walter Scheithauer
1946 Born in Germany.
1969 B. A. California State University.
1973 M. D. Loma Linda University, CA.
1974~1978 Resident, Stanford University, CA.
1979~1980 Instructor, Pathology, Mayo Medical School.
1980~1984 Assistant Professor.
1980~1987 Associate Professor.
1987~ Professor, Pathology, Mayo Medical School.
of ML was well known as "ghost tumor", described by Vaquero et al. in 1984. Radiation and following COP chemotherapy were done, although repeated warnings were given by the pathological side with a kind advice of Prof. Scheithauer (demyelinating disease: 1991. 6. 12.). She was doing well till the present recurrence (disorientation as to time, short concentration span, untidy manner, ataxia in the lt. extremities and unsteady gait with the tendency to fall toward the left. Recent CT and MRI showed multifocal enhanced foci in rather selective involvement of white matter in rt. basal ganglia, lt. periventricular white matter, splenium of corpus callosum and both middle cerebellar peduncles (Fig. 1). For sure diagnosis, CT—guided stereotactic biopsy was performed on 9. Apr. 1992.

Histologically there were multiple infiltration foci of large lymphoid tumor cells, resembling the tumor cells in the lymph node specimen that had been extirpated in 1982. Many mitotic figures were seen (Fig. 2). The immunostaining showed abundant expression of the B—cell marker (L–26) in the cell membrane and the cytoplasm of many tumor cells (Fig. 3). Tumor cells were stained also for leukocyte—commonantigen (Fig. 4), but never developed T—cell markers (MT—1 or UCHL).

Comments: Dr. Scheithauer pointed out that this case was a typical pitfall on the diagnosis of malignant lymphoma. He was not so informed of the clinical data, especially about preoperative administration of high dose of dexamethasone.

The instructive fact was that first we should not use glucocorticoids prior to biopsy on suspected
malignant lymphoma. Second stereotactic biopsy was very useful for the diagnosis of malignant lymphoma as shown by Dr. Scheithauer et al.

References:

Intracranial Plasmacytoma

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[Case]
A 48-year-old man was admitted to our hospital with slowly progressive gait disturbance. On admission, neurological examination revealed mild right hemiparesis. Plain X—ray films of the skull were normal. The CT scan showed an irregular enhanced lesion with severe edema in the left frontal lobe. The MR image revealed the lesion located at the medial part of left frontal cortex with spreading into interhemispheric fissure and brain surface diffusely.

At operation, the elastic hard greyish tumor like lesion was found in the moderately edematous left frontal cortex. The dura matter, Falx cerebri, and arachnoid membrane were thickened at this portion. Subtotal removal of tumor was performed.

[Pathology]
Histologically, the lesion consists of fibrovascular tissue with heavy infiltrates of plasma cells. Scattered lymphocytes (B—cell) were also present. Whorl formations and psammomatous bodies were absent. The PAS and Grocott stains revealed no organisms.

[Comments]
With marked plasma cell infiltrate, plasmacytoma was considered. In this case, immunohistochemical analysis demonstrated the polyclonal nature of the plasma cell proliferation. From this point of view, it may be inflammatory proliferation. (e. g. plasma cell granuloma)

Does true polar spongioblastoma exist?

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This 45 y—o, right—handed housewife (T. O., 268692) was admitted to our ward complaining of unconscious attack of late onset. On admission neurological exam. revealed no abnormality. Her family and past histories were unremarkable. A CT scan showed an isodensity round mass with slight edema in left frontal lobe. (Fig. 1) The tumor had homogenous contrast enhancement. Angiography disclosed an avascular mass with round shift of ACA to the left. At operation, grayish, soft tumor
This 45 y-o female had two episode of unconscious attack three years before admission. On CT scan, round, well circumscribed, isodensity mass with slight edema was recognized (upper) and homogenously enhanced (lower).

The tumor specimen shows highly distinctive cytoarchitecture like “terraced field”. Cells are arranged in parallel fashion of 2–3 rows between acellular zone. In some areas the feature of oligodendroglioma such as “fried egg” was recognized among vascular connective tissue stroma. (inset) H & E ×100 & 200

There are some strong GFAP reactivity in tumor cells. GFAP stain ×200
was well—demarcated and removed en bloc. Post-operative course was uneventful and the patient received irradiation and chemotherapy. Now over 7 years after operation, she is still living, somewhat obtunded, probably due to radiation therapy and tumor—free on follow—up CT scans.

On histological exam. the tumor specimen showed highly distinctive arrangement of cells like “terraced field”. (Fig. 2) The cells were arranged in parallel fashion of 2–3 rows, forming fairly compact bands or palisades. The tumor cell groups were separated by a delicate and regularly arranged vascular connective tissue stroma. Typical cell arrangement of oligodendroglioma and minimum anaplastic changes such as small necrosis were recognized in some area (Fig. 2 inset). On GFAP immunostain, some were positive and some were negative. (Fig. 3) GFAP—positive cells were intermingled and this showed that positive cells were not reactive but tumor cells. Our histological diagnosis was mixed oligo—astrocytoma. Prof. Scheithauer diagnosed this fascinating tumor as polar spongioblastoma and said that the tumor should not be called a mixed lesion but polar spongioblastoma with focal oligodendrogial differentiation. The true polar spongioblastoma as originally defined by Russel and Cairns in 1947 seen in the preparations of the cerebrum in 16—week—old embryo. On neuropathological practice, however, we have rarely experienced those typical polar spongioblastoma as well as astroblastoma. Dr. Rubinstein described that occasionally foci of “pure” polar spongioblastoma may, like astroblastic foci, be met in isolated fields in glioblastoma or dedifferentiating diffuse cerebral astrocytomas. We performed A2B5 immunostain. A2B5mAb was purchased from ATCC (American Type Culture Collection) and this stains type 2 astrocytes and oligodendrocyte—type 2 astrocyte progenitor cells. We found that there were weak reaction as seen in mixed gliomas. Further detailed study whether true polar spongioblastoma exist or not should be made.

Acute desseminated encephalomyelitis (ADEM) mimicking a temporal lobe glioma

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This 19—year—old male was well until January 14th, 1992, when he developed headache and visual field defect. Neurological examination was unremarkable except for complete right homonymous hemianopsia. He was afebrile. CT scan showed left temporal low density area partially enhanced by contrast material (Fig. 1). MR imaging revealed left temporal swelling and loss of normal gray and white matter structures. The lesion was heterogeneously enhanced by Gadolinium—DTPA. There was no other detectable lesion. On the day of admission, he had a general convulsion. Operative findings (01—29—92): Corticotomy revealed grayish elastic hard tumor in the left temporal lobe. Frozen section of the biopsy specimen was reported as a low grade astrocytoma. The temporal lobectomy was performed.

Postoperative course: His immediate postoperative course was uneventful but a week later, he...
had fever and persistent hiccup, followed by obtundation and respiratory distress which required intubation and respirator. Repeated MRI showed extensive lesions in the midbrain, pons, medulla and cerebellum. Lumbar CSF contained 388 WBCs (Poly 60%, Lymph 40%), total protein 99mg/dl, and glucose 55mg/dl. CSF cytology revealed moderate number of atypical immature lymphocytes. Both CSF oligoclonal band and myelin basic protein were negative. Extensive virological studies, including EB, HS, HZ, rubella, and mumps except for rubeola (1 : 16) were all negative. Daily Aciclovir injection did not alter the clinical course but steroid pulse therapy was very effective. He recovered well and his right homonymous hemianopsia improved. Repeated MRI showed resolution of the brain stem lesions.

**Histological findings:** The lesion contains foci of necrosis and hemorrhage with prominent perivascular lymphocytic cuffings (Fig. 2). These lymphocytes are mixture of T¬ and B¬-cells and there are areas of reactive astrocytosis and demyelination.

**Comments:** With the advent of MR imaging, there are increasing case reports of ADEM but its histological findings are rarely reported. ADEM usually manifests itself as multiple CNS lesions but when initial radiological studies reveal a single lesion, diagnosis of ADEM is difficult without a biopsy. Although it is rare, ADEM should be added to the list of the differential diagnosis for intraparenchymal lesion.

**A recurrent case of hemangioblastoma**

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**Case:** T. A., a 34—year old male

**CC:** Gait disturbance and speech disturbance

**PI:**

On April 9, 1986, the patient underwent resection of right cerebellar hemangioblastoma at a medical university. He developed right hearing disturbance, cerebellar ataxia, and right facial palsy after operation, but these symptoms were considerably improved.

Around the middle of January 1992 his speech became disconnective and his gait was rather unsteady. Headaches also became worse. He consulted our clinic in February 1992. Since performed CT scans revealed abnormality, he was hospitalized in our hospital on March 9, 1992.

**Neurological examinations:**

There were ataxic gait, speech disturbance, right
hearing disturbance, and right facial palsy.

CT and MRI findings:

There was a solid mass in the center of the right cerebellopontine angle region with a cyst in the medial part, which compressed the brain stem. Contrast enhancement clearly revealed a solid mass. (Fig. 1)

Operation:

On March 19, 1992 subtotal removal of the right cerebellar tumor was performed through the right suboccipital craniectomy.

A soft hemorrhagic solid tumor and yellowish cyst existing in the right cerebellopontine angle to inside the cerebellum were resected. The exterior part of the tumor was rather soft with relatively clear margin.

Histology:

Some parts of the tumor show a typical hemangioblastoma. Another parts of the tumor show slender processes with positive GFAP staining and many endothelial proliferation. (Fig. 2)

Diagnostic problem:

Recurrence of hemangioblastoma with gliosis or glioma.

Comments:

Prof. Scheithauer suggested that the tumor recurred undoubtedly as a hemangioblastoma and malignant glioma might be induced with a hemangioblastoma by irradiation.