8-14 translocation in a Japanese Burkitt’s lymphoma.

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Abstract

Chromosome analysis was performed on cells obtained from the pleural effusion of a Japanese patient with Burkitt’s lymphoma. Two modal chromosomal numbers were found: 45 and 46. Five different karyotypes were present, all having a t (8q−;14q+) translocation. This case illustrates that Burkitt’s lymphomas of Japanese are no exception to the frequent association of this chromosomal abnormality with Burkitt’s lymphomas.

KEYWORDS: Japanese Burkitt’s lymphoma, 8-14 translocation, chromosome analysis.

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8-14 TRANSLOCATION IN A JAPANESE BURKITT'S LYMPHOMA

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Abstract. Chromosome analysis was performed on cells obtained from the pleural effusion of a Japanese patient with Burkitt's lymphoma. Two modal chromosomal numbers were found: 45 and 46. Five different karyotypes were present, all having a t(8q−;14q+) translocation. This case illustrates that Burkitt's lymphomas of Japanese are no exception to the frequent association of this chromosomal abnormality with Burkitt's lymphomas.

Key words: Japanese Burkitt's lymphoma, 8-14 translocation, chromosome analysis.

Cytogenetic studies on African, North American and European Burkitt's lymphomas disclosed the presence of a specific t(8q−;14q+) translocation (1-4). Recently, we reported on the Burkitt's lymphoma of a Japanese, in which a 14q+ marker chromosome was found but the origin of the extra chromosomal material attached distally to the long arm of 14 chromosome could not be determined (5). We describe here the t(8q−;14q+) translocation in another Japanese patient with Epstein-Barr virus negative Burkitt's lymphoma.

A 17-year-old boy was in good health until mid-May 1979, when he developed abdominal distention. This was followed in mid-June by fatigue, anorexia and vomiting. Abdominal masses were palpated by his family physician. A gastrointestinal series demonstrated multiple intra-abdominal tumors displacing the intestinal loops. The patient was referred to Okayama University Hospital, where an exploratory laparotomy on July 17 revealed a massive tumor involving the retroperitoneum and mesentery and an unresectable intestinal narrowing. Biopsy material tumor was not submitted for pathological examination due to a mishap. However, surface marker analysis of the tumor cell suspension indicated 90% surface immunoglobulinpositive cells. There was no lymph-
adenopathy or hepatosplenomegaly. Laboratory tests on admission were all normal. Several courses of chemotherapy with various combinations of drugs induced partial tumor regression each time, but tumor regrowth was rapid. Eventually, the disease became refractory to chemotherapy and the patient died of intestinal obstruction on December 25, 1979. Serum antibody titers against Epstein-Barr virus capsid antigen and nuclear antigen were 1:40 and 1:80, respectively, but antibodies against early antigen were not detectable at a titer of 1:10. Tumor cells from the pleural effusion were negative for EBNA. Autopsy material from the abdominal tumor revealed undifferentiated lymphoma with a starry sky pattern.

Chromosomes were analyzed on tumor cells from the pleural effusion aspirated four days prior to death. The cells were separated on a Ficoll-Conray gradient, incubated at 37°C for 15 h in McCoy 5a medium containing 20% fetal calf serum and then treated for two h with Colcemid (0.5 μg/ml). Cells in metaphase were swollen with 0.075 M KCl for 13 min at 37°C and fixed in a methyl alcohol : acetic acid (3:1) solution. Chromosome preparations were air-dried and banded with quinacrine mustard (6).

As shown in Table 1, two modal chromosomal numbers, 45 and 46, were found. Variations found in the karyotyped cells are summarized in Table 2.

**Table 1. Chromosome number distribution found in tumor cells from pleural effusion**

<table>
<thead>
<tr>
<th>Chromosome number</th>
<th>45</th>
<th>46</th>
<th>90</th>
<th>92</th>
<th>Total</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>19 (3)</td>
<td>29 (8)</td>
<td>1</td>
<td>1</td>
<td>50 (11)</td>
</tr>
</tbody>
</table>

Number in parentheses = number of banded cells examined.

**Table 2. Variation found in the karyotyped cells from pleural effusion**

<table>
<thead>
<tr>
<th>Variation</th>
<th>46, XY, t[8;14] (q24;q32), 17p−</th>
<th>46, Xp+, Yq+, t[8;14] (q24;q32), 17q+, 17p−</th>
<th>45, XY, t[8;14] (q24;q32), 17p−, 18q−, 18q+</th>
<th>45, XY, t[8;14] (q24;q32), −13, 17p−</th>
<th>45, X, −Y, t[8;14] (q24;q32), 17p−</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>5</td>
<td>3</td>
<td>1</td>
<td>1</td>
<td>1</td>
</tr>
</tbody>
</table>

Five different karyotypes were present, all having a t(8q−;14q+) translocation. Of the 11 karyotyped metaphase cells, 5 showed a karyotype of 46, XY, t[8;14] (q24;q32), 17p−, 3 a karyotype of 46, Xp+, Yq+, t[8;14] (q24;q32), 17q+, 17p−, (Fig. 1) and the remaining 3 each a different karyotype. Extra chromosomal bands were observed on the short arm of chromosome X (Xp+)
and on the long arms chromosomes 17, 18 and Y (17q+, 18q+ and Yq+) but their origin could not be determined. The short arm of chromosome 17 appeared to have a deletion at band 17p11.

In the present study, we have described a typical t(8q-; 14q+) translocation in a Japanese boy with Burkitt's lymphoma. The same translocation has recently been detected in a 6-year-old Japanese boy with Burkitt's lymphoma.
admitted to the Department of Pediatrics (personal communication, 1979 by Kei Nagase). These two cases along with the one previously reported by us (5) strongly suggest that the 14q+ abnormality may be a frequent occurrence in Burkitt's lymphomas of Japanese as with those of African, North American, and European origin.

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REFERENCES