Diagnostic relevance of peripheral blood immunocytochemistry in hairy cell leukaemia

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Abstract

Aims—(1) To assess the diagnostic relevance of peripheral blood immunocytochemistry in hairy cell leukaemia (HCL); (2) to compare the immunostaining of bone marrow biopsy specimens with bone marrow and peripheral blood cytospins; (3) to evaluate the sensitivity of the different markers used; (4) to identify the ultrastructural localisation of DBA.44 in HCL variant.

Methods—Immunoenzymatic staining procedures, immunoperoxidase and immunoperoxidase alkaline phosphatase, were used with a panel of monoclonal antibodies directed to HCL associated antigens. Ultrastructural immunostaining was performed using colloidal gold conjugated antibodies.

Results—HCL showed strong cytoplasmic reactivity for CD22, CD25, CD103, DBA.44, κ, or λ light chains. Peripheral blood diagnostic hairy cells were found in all the cases with absolute counts ranging from 0.11 × 10⁹/l up to 6.4 × 10⁹/l and values increasing with the size of the spleen. A median of 36.5% of leukaemic cells was found in bone marrow aspirates and 70% in bone marrow trephine specimens. The monoclonal antibodies CD22 and DBA.44 showed the highest and the lowest percentage of positive hairy cells, respectively; this difference was statistically significant (p = 0.0025). Ultrastructural immunolabelling with DBA.44 showed a cytoplasmic membrane localisation of the antigen in one case of HCL variant.

Conclusions—(1) Immunocytochemistry is a useful technique which enhances the accuracy of diagnosis in HCL; (2) peripheral blood immunocytochemistry is recommended because it highlights hairy cells in all cases; (3) CD22 appears to be the most sensitive of the markers tested; (4) ultrastructural analysis is a useful tool in selected cases of HCL variant.

Keywords: Immunocytochemistry, hairy cell leukaemia, diagnosis.

Hairy cell leukaemia (HCL) is a chronic B cell lymphoproliferative disorder which accounts for 2% of all forms of leukaemia in adults. It is characterised by splenomegaly, pancytopenia, bone marrow infiltration, and circulating leukaemic cells with typical cytoplasmic projections. Recent studies have shown that HCL has a distinctive, unique pattern of antigen expression.

Bone marrow aspirates are often unsuccessful in HCL (dry tap) because of a marked increase in reticulin fibres. Diagnostic material is obtained by aspiration only in a minority of patients and hairy cells are often undetectable in conventionally stained peripheral blood films because of the pancytopenia. Bone marrow trephine is considered the most reliable diagnostic tool for HCL, and immunohistochemistry can help when the presence of an infiltrate is considered equivocal by standard microscopy.

Immunocytochemistry on bone marrow and peripheral blood cytospins could be a useful tool for the detection of hairy cells because morphology and immunophenotype determination can be combined. The aims of this study were to assess the diagnostic relevance of peripheral blood immunocytochemistry in HCL, to compare peripheral blood with bone marrow immunostaining, and to evaluate the sensitivity of the different reagents used. Bone marrow trephine biopsy specimens and bone marrow and peripheral blood cytospins from 23 HCL patients were studied at diagnosis using a panel of monoclonal antibodies directed to HCL associated antigens. As ultrastructural analysis of the cell can be relevant in the differential diagnosis of closely related types of chronic B cell leukaemia, ultrastructural examination of peripheral blood was conducted in two cases of HCL variant to provide a better morphological evaluation of the leukaemic cells and to identify the localisation of the HCL associated antigen detected by the monoclonal antibody DBA.44.

Methods

MATERIALS

Twenty one patients with HCL and two with HCL variant were studied at diagnosis. All subjects included in this study were referred to our institution between July 1992 and June 1994. Nineteen of the patients with classic HCL were male and two were female. Median age was 54 years (range 38–72). Median haemoglobin value was 12.4 g/dl (range 7.6–15.2). The white blood count varied from 1.0 × 10⁹/l to 11.2 × 10⁹/l (median 2.5 × 10⁹/l). The absolute platelet counts ranged from 20 × 10⁹/l to 138 × 10⁹/l (median 72 × 10⁹/l). According to the spleen size, patients were subdivided for analysis into three groups: (1) patients with no palpable spleen (n = 4), (2) patients with splenomegaly <5 cm below the
Table 1  Antibody reagents used in this study

<table>
<thead>
<tr>
<th>Antibody</th>
<th>CD No.</th>
<th>Specificity</th>
<th>Source</th>
</tr>
</thead>
<tbody>
<tr>
<td>OKB-22</td>
<td>CD22</td>
<td>Pan B cell</td>
<td>Ortho</td>
</tr>
<tr>
<td>L26</td>
<td>CD26</td>
<td>B cell</td>
<td>Dakopatts</td>
</tr>
<tr>
<td>B-ly-7</td>
<td>CD103</td>
<td>HCL T cell subset</td>
<td>Dr 3 Poppena (Canada)</td>
</tr>
<tr>
<td>DBA-44</td>
<td>Non-clustered</td>
<td>HCL B cell subset</td>
<td>Dakopatts</td>
</tr>
<tr>
<td>ACT-1</td>
<td>CD25</td>
<td>IL-2 Rec a chain</td>
<td>Dakopatts</td>
</tr>
<tr>
<td>UCHT-1</td>
<td>CD3</td>
<td>Pan T cell</td>
<td>Dakopatts</td>
</tr>
<tr>
<td>Lysozyme</td>
<td>Non-clustered</td>
<td>Monocytes</td>
<td>Dakopatts</td>
</tr>
<tr>
<td>kappa</td>
<td>Non-clustered</td>
<td>Cycling cells</td>
<td>Dakopatts</td>
</tr>
<tr>
<td>lambda</td>
<td></td>
<td></td>
<td>Dakopatts</td>
</tr>
</tbody>
</table>

costal margin (n = 9), (3) patients with splenomegaly (>5 cm below the costal margin (n = 8).

The two patients with HCL variant showed marked splenomegaly (>5 cm below the costal margin) and white blood cell counts of 41 x 10^6 and 11 x 10^6 cells/l, respectively.

Normal control bone marrow and peripheral blood cells were obtained from five healthy adults and control bone marrow core biopsy specimens from five patients with normal cellularity. Informed consent for using cells in this study was obtained from all the subjects. The diagnosis of HCL was based on clinical features, cell morphology, immunophenotype, histology, and, in the two cases of HCL variant, by ultrastructural analysis.

PROCEDURES

Mononuclear cells were isolated from heparinised peripheral blood (n = 23) and bone marrow samples (n = 15) by Ficoll-Hypaque density gradient centrifugation. Cytospins were prepared with a concentration of 5 x 10^5 cells per slide, air dried overnight, wrapped in aluminium foil, and stored at -20°C until immunostained. Immunocytochemical labelling was performed by the alkaline phosphatase-antialkaline phosphatase (APAAP) and immunoperoxidase (IP) techniques, as previously described. 10

The antibody reagents used are listed in table 1. The proportion of positive cells was evaluated by light microscopy with oil immersion (magnification x 1000), examining 1000 lymphoid cells per sample. Only cells showing unequivocal morphological appearance of hairy cells (large size, eccentric nucleus, abundant cytoplasm, and characteristic long cytoplasmic villi) were considered to be leukemic.

Bone marrow core biopsy specimens (n = 23) were fixed in 10% buffered formalin, decalcified in Decal, and embedded in paraffin wax. Five micron thick sections were cut from paraffin wax blocks and stained with Giemsa and Gomori silver impregnation for reticulin fibres. Immunohistochmical studies were performed on paraffin wax sections by the APAAP method11 using the monoclonal antibodies L26 (CD20) and DBA.44. Slides were examined by light microscopy and the percentage, pattern of distribution, and morphological characteristics of positive cells were evaluated.

The ultrastructure of two HCL variant cases was examined on peripheral blood mononuclear cells isolated by Ficoll-Hypaque density gradient centrifugation.

For morphological evaluation, the samples were fixed in 2.5% buffered glutaraldehyde for 1 h, postfixed in 1% osmium tetroxide (OsO₄), and embedded in Epon.

For ultrastructural immunostaining, the cells were fixed in 4% phosphate buffered formaldehyde for 10 minutes, permeabilised with Triton X-100 (0.25% in phosphate buffered saline (PBS)) for 5 minutes, and incubated for one hour with CD22 and DBA.44 monoclonal antibodies (diluted 1:5 in PBS). After a second fixation in 2.5% buffered glutaraldehyde for 10 minutes, samples were incubated for one hour with a 10 nm colloidal gold conjugated goat anti-mouse antibody (BioCell Research Laboratories) diluted 1:10 in PBS, postfixed in 1% OsO₄, and embedded in Epon. All the incubations were performed at room temperature. Semithin and ultrathin sections were stained with toluidine blue and uranyl acetate, respectively, and observed with a Philips CM10 electron microscope.

Comparison between groups were analysed using the Student t test; linear regression analysis was carried out to study the possibility of a trend across ordered groups.

RESULTS

EXPRESSION OF HCL ASSOCIATED ANTIGENS IN NORMAL BONE MARROW AND PERIPHERAL BLOOD CELLS

Immunocytochemical staining of cytopsins from five normal peripheral donors showed a median of 0.3% (range 0.1-0.7) CD103+ cells and 2.2% (range 2.1-2.7) DBA-44+ cells in the bone marrow, and 0.6% (range 0.2-0.9) CD103+ and 4.2% (range 3.2-5.5) DBA-44+ cells in the peripheral blood respectively. These were small round lymphoid cells. DBA-44 highlighted short thin cytoplasmic projections in a minority of peripheral blood small lymphocytes. No CD22+, CD103+, DBA.44+, or CD25+ cells with morphology of hairy cells were found in the bone marrow or peripheral blood of normal individuals.

Bone marrow biopsy specimens from five patients with normal cellularity showed rare scattered DBA.44 positive small lymphocytes. The staining pattern was membrane associated; no cytoplasmic projections were seen.

IMMUNOCYTOCHEMICAL Staining of peripheral blood and bone marrow cytopsins in HCL

HCL showed a very consistent immunophenotype. Strong cytoplasmic reactivity of hairy cells for CD22, CD25, CD103, DBA.44, κ, or λ light chains was found in bone marrow and peripheral blood cytopsins from the 21 cases of HCL (table 2). Labelled hairy cells were clearly distinguished from non-leukaemic cells and the typical morphology was well preserved (fig 1).

Peripheral blood circulating hairy cells were easily identified by immunocytochemistry in all the cases. Seventy six percent of the patients had >10% peripheral blood hairy cells and only one patient had less than 2% detectable hairy
Table 2  Percentage of positive cells on cytopsin slides. Values are means (SD)

<table>
<thead>
<tr>
<th>Antibody</th>
<th>BM aspirate (n=14)</th>
<th>Ph (n=21)</th>
<th>Cell type</th>
</tr>
</thead>
<tbody>
<tr>
<td>CD22</td>
<td>40 (18)</td>
<td>31 (29)</td>
<td>Hairy cells</td>
</tr>
<tr>
<td>CD103</td>
<td>35 (19)</td>
<td>26 (28)</td>
<td>Hairy cells</td>
</tr>
<tr>
<td>DBA-44</td>
<td>26 (16)</td>
<td>22 (17)</td>
<td>Hairy cells</td>
</tr>
<tr>
<td>CD25</td>
<td>38 (18)</td>
<td>28 (27)</td>
<td>Hairy cells</td>
</tr>
<tr>
<td>kappa</td>
<td>25 (28)</td>
<td>10 (17)</td>
<td>Hairy cells</td>
</tr>
<tr>
<td>lambda</td>
<td>16 (20)</td>
<td>20 (30)</td>
<td>Hairy cells</td>
</tr>
<tr>
<td>CD3</td>
<td>31 (15)</td>
<td>62 (26)</td>
<td>T lymphocytes</td>
</tr>
<tr>
<td>α-Lys</td>
<td>Not done</td>
<td>1-4 (1-3)</td>
<td>Monocytes</td>
</tr>
</tbody>
</table>

Hairy cells. The proportion of circulating CD22 positive hairy cells varied from 1-32% to 98% of mononuclear cells (median 20%) with absolute count ranging from 0-11 x 10^9/l up to 6-4 x 10^9/l. Among the monoclonal antibodies directed against HCL associated antigens, CD22 accounted for the highest [31(SD 29)%] and DBA.44 for the lowest [22(25)%] percentage of positive hairy cells (table 2). The difference of sensitivity between CD22 and DBA.44 was statistically significant (p=0.0025). The pattern of immunostaining with DBA.44 showed more intense membrane staining than with CD22. The characteristic projections of the leukaemic cells were strongly highlighted by both antibodies. The percentage of CD103 [26(28)%] and CD25 [28(27)%] positive hairy cells was lower than with CD22; however, this difference was not significant. Monoclonal light chain was identified in all patients, including those with a small peripheral blood infiltration. Thirteen cases were λ light chain positive (62%) and eight were κ light chain positive. The typical monocytopenia, with a median of 1% (range 0-1-6%) antilysozyme positive monocytes was observed in all cases. A negative correlation was found between the percentage of peripheral blood hairy cells and the proportion of CD3 positive T lymphocytes (table 2).

Bone marrow aspirate was unsuccessful (dry tap) in one case and markedly hypocellular in two cases. Diagnostic material was available in 14/15 patients. A proportion of >10% hairy cells was found in all the bone marrow samples. The percentage of CD22 + hairy cells in bone marrow ranged from 16% to 69% (median 36-5%). Similar results were shown with CD103 and CD25. In contrast, DBA.44 accounted for a significant lower percentage of positive hairy cells (table 2).

Figure 1  Peripheral blood cytopsin immunostaining with APAAP (red cytoplasmic staining) shows two CD22 positive hairy cells with eccentric nucleus and abundant cytoplasm with characteristic vili, and a small CD22 positive B lymphocyte.

Table 3  Evaluation of cellularity, L26, DBA.44, fibrosis, and pattern of infiltration in bone marrow biopsy specimens of 21 HCL patients

| Cellularity (%) | 70 | 80 | 60 | 60 | 70 | 20 | 40 | 90 | 25 | 60 | 60 | 80 | 40 | 50 | 70 | 50 | 70 | 60 | 50 | 25 | 40 |
|-----------------|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|----|
| L26(%)          | 50 | 50 | 50 | 50 | 50 | 50 | 50 | 70 | 65 | 65 | 80 | 80 | 80 | 50 | 50 | 80 | 65 | 65 | 80 | 80 | 50 | 45 | 80 |
| DBA-44(%)       | 5  | nd | 5  | 20 | 40 | 10 | 25 | 30 | nd | 20 | nd | 50 | nd | 40 | 20 | 25 | 65 | 5  | 25 | 25 | 10 |
| Fibrosis Pattern| I = interstitial; D = diffuse; nd = not done
| Pattern         | I = interstitial; D = diffuse; nd = not done

A median of 26% (range 5-65%) DBA.44 + leukaemic cells was observed, a proportion significantly lower to the percentage of L26 + cells (table 3). Immunohistochemical labelling gave a more accurate assessment of hairy cells than morphology alone. In two cases with a low degree of interstitial bone marrow infiltration, immunohistochemistry permitted easier identification of the leukaemic population than did morphology. In one case with marked fibrosis, hypoplasia, and scattered CD20 and DBA.44 positive hairy cells in the trephine sample, immunocytochemical staining of bone marrow and peripheral blood cytopsin—with 16% and 5% hairy cells respectively—was very helpful in confirming the diagnosis of HCL.

Neither the amount of bone marrow infiltration assessed by L26 positivity in trephine biopsy specimens nor bone marrow fibrosis was correlated with the percentage or absolute number of peripheral blood leukaemic cells assessed by CD22 (>0.1). The percentage and absolute number of hairy cells in peripheral blood was found to increase with spleen size. In contrast, no correlation was found between
Figure 2 Bone marrow trephine and peripheral blood infiltration by leukaemic cells according to the spleen size. Values represent the mean of the percentage of CD22+ peripheral blood hairy cells and L26+ bone marrow hairy cells (A), and of the absolute number of peripheral blood hairy cells (B) in patients subdivided according to the spleen size.

the size of the spleen and the proportion of bone marrow hairy cells (fig 2).

MORPHOLOGY, IMMUNOPHENOTYPE, AND BONE MARROW TREPHINE OF HCL VARIANT Light microscopic examination of peripheral blood showed a homogeneous population of cells with round nucleus, condensed chromatin, and prominent nucleoli. In one case a prominent nucleolus was present in only a minority of the cells. The cytoplasm was slightly basophilic with fine hairy projections. The proportion of circulating CD22+ hairy cells was 99% (34.4 × 10⁹/l) and 89% (6 × 10⁹/l) in the two patients. Leukaemic cells were CD22 and DBA.44 positive but CD25 and CD103 negative. Compared with CD22, DBA.44 showed more intense cytoplasmic membrane staining. The percentage of cells in cell cycle (Ki67+) was 3.9% and 0.5% respectively. Bone marrow trephine biopsy specimens showed minimal involvement by leukaemic cells (20% of the total cellularity). The pattern of infiltration was interstitial, with leukaemic cells within the lumen of bone marrow capillaries; reticulin fibres were slightly to moderately increased.

ELECTRON MICROSCOPY OF HCL VARIANT Ultrastructural examination of the two HCL variant cases showed medium sized cells with round nuclei and heterochromatin condensation at the nuclear periphery. A prominent nucleolus was observed in both cases. The cytoplasm contained numerous mitochondria and free polyribosomes, with strands of long rough endoplasmic reticulum characteristically circumscribing the nucleus. All leukaemic cells showed thin cytoplasmic projections as well as broad based and large buds with an uneven distribution around the membrane. No ribosome lamellar complexes were observed.

Ultrastructural immunolabelling with CD22 and DBA.44 monoclonal antibodies showed gold particles staining leukaemic cells. Immunolocalisation appeared to be precise and background signal was low. Colloidal gold particles corresponding to CD22 uniformly covered the cell membrane and were also observed in the cytoplasm, apparently not related to any specific structure. By contrast, colloidal gold particles corresponding to DBA.44 were only observed on the cell membrane and were more heavily distributed along the villous projections of the cytoplasm (fig 3A and B).

Discussion

The accurate diagnosis of hairy cell leukaemia has become important after the development of highly effective therapeutic agents (α-interferon, deoxycoformycin, 2-chlorodeoxyadenosine) that can induce responses in a high proportion of patients.¹²⁻¹⁴ The simultaneous morphological and immunological evaluation of hairy cells is a powerful tool for the diagnosis of HCL. Immunostaining with monoclonal antibodies is a valid adjunct to conventional morphology in the identification of leukaemic cells in HCL.³⁻⁵,¹⁵,¹⁶ Using immunocytochemistry with a combination of antibodies, we regularly identified as few as 1% hairy cells in peripheral blood and bone marrow samples.

Among the monoclonal antibodies proposed as useful tools for the diagnosis of HCL,²⁻⁶ we excluded CD11c and HC2 because both strongly react with monocytes and myeloid cells which can be morphologically similar to hairy cells. In our hands, the CD22 monoclonal antibody appears to be the most sensitive marker for the diagnosis of HCL. Unlike the other HCL associated antigens, the majority of hairy cells were CD22 positive and the APAAP staining makes CD22 positive hairy cells easily distinguishable from the small positive normal lymphocytes because of their larger size and cytoplasmic projections.

We also compared immunostaining of hairy cells from different sources. Immunocytochemical staining of peripheral blood cytospins allowed the identification of leukaemic hairy cells, usually overlooked by morphological examination, in all the cases and proved to be helpful in supporting the diagnosis in those patients with minimal bone marrow infiltration. The absolute number of hairy cells was variable ranging from 0.1 × 10⁹/l to 6.4 × 10⁹/l; however, the phenotype was easily characterised, even in those patients with small numbers of circulating leukaemic cells.

The percentage and absolute number of peripheral blood leukaemic cells increased with
Diagnostic immunocytochemistry in hairy cell leukaemia

These monoclonal antibodies, highlighting the characteristic cytoplasmic projections, were also able to detect scattered individual leukemic cells in hypoplastic bone marrow. In contrast to a previous report, the sensitivity of DBA.44 was significantly lower than that of the L26 monoclonal antibodies. Because the small numbers of cells were enough to make cytopsins, bone marrow aspiration provided sufficient diagnostic material in 14/15 patients, with a proportion of diagnostic hairy cells greater than 10% in all the samples. We found a lower percentage of hairy cells in bone marrow cytopsins than in bone marrow trephine specimens; the high proportion of CD3 positive lymphocytes on bone marrow cytopsins, however, indicates that this difference was due to peripheral blood contamination.

Although HCL is now well established as a B cell leukaemia, its normal counterpart in B cell ontogeny remains undefined. Most investigators agree that HCL is at a late stage of B cell differentiation. As previously reported, we have found a subpopulation of CD103+ lymphocytes in normal peripheral blood. Moreover we have identified a normal peripheral blood subpopulation of DBA.44+ lymphocytes ranging from 3-2% to 5-5% of the mononuclear fraction, a minority of which show thin cytoplasmic projections. If a normal counterpart of the hairy cells exists, it may include a subset of DBA-44+ cells.

Immunocytochemistry seems to be promising to monitor treated patients for residual disease after chemotherapy. Using the combination of markers described in this study, we have detected minimal residual disease in bone marrow and peripheral blood of HCL patients considered to be in complete remission by conventional criteria of response to treatment. DBA.44 has been described as being helpful in the monitoring of residual disease. This study shows that CD22, CD25, and CD103 monoclonal antibodies have higher sensitivity than DBA.44 in identifying leukaemic hairy cells at diagnosis. Moreover, we found the same difference in sensitivity in the evaluation of minimal residual disease (manuscript in preparation).

HCL variant is an uncommon disorder. We found two HCL variant cases out of 23 consecutive HCL patients referred to our institute in a two year period. The distinction of HCL variant from other lymphoproliferative disorders such as splenic lymphoma with villous lymphocytes (SLVL) and prolymphocytic leukaemia (PLL) may be difficult. Defining criteria of these disorders are based on the appraisal of the relevant morphological features as diagnostic combinations of immunological markers, as described for classical HCL, have not been identified.

We found immunocytochemistry to be helpful in the diagnosis of HCL variant. The combination of hairy morphology, a high proportion of peripheral blood infiltration (90% leukemic cells), and the lack of some of the “HCL associated” antigens (CD25 and CD103 negative), led us to consider the diagnosis of HCL

spleen size. The small number of patients does not permit any statistical evaluation; the trend, however, supports the hypothesis that the circulating hairy cells originate from the spleen and spread to the peripheral blood.

The antilysozyme antibody was also helpful in the diagnosis of HCL by highlighting the low percentage of monocytes (1%), in patients with low peripheral blood leukemic infiltration.

The bone marrow trephine biopsy is considered the most reliable diagnostic tool for HCL because of the high rate of “dry tap” caused by fibrosis. On the bone marrow trephine specimens, L26 and DBA.44 were useful in confirming the diagnosis of HCL and estimating the degree of bone marrow infiltration.

Figure 3 (A) Ultrastructural immunolabelling of HCL variant shows colloidal gold particles (black dots) corresponding to DBA.44 monoclonal antibodies uniformly decorating the cytoplasmic membrane, with a more heavy distribution along the villous projections of the cytoplasm; × 8140. (B) Colloidal gold particles corresponding to DBA.44 (black dots) covering the cytoplasmic villi; × 19 350.
variant. It is worthwhile noting that, despite the high white blood cell count, the percentage of cells in cell cycle (Ki67+) was low.

Morphological evaluation of the leukaemic cells by electron microscopy highlighted prominent nucleoli in both HCL variant cases. This supported the diagnosis of HCL variant in the case where prominent nucleoli were observed by light microscopy only in a minority of the leukaemic cells. Ultrastructural immunostaining with DBA.44 in HCL variant showed membrane localisation of this antigen. This could explain its intense cytoplasmic membrane staining by immunocytochemistry.

Our results are in agreement with a larger series.5 However this is the first study that compared bone marrow, peripheral blood cytosin, and bone marrow trephine specimen immunostaining in the evaluation of the sensitivity of the markers used. We conclude that immunocytochemistry is a useful method which, in combination with routine morphology and histology, significantly enhances the accuracy in diagnosis of HCL and HCL variant. CD22 appears to be a valuable marker, but the application of a complete antibody panel is still recommended for higher specificity and sensitivity in the diagnostic procedure. Peripheral blood immunocytochemistry is always recommended, being diagnostic (HC>10%) in the majority of cases; long term follow up will establish whether the proportion of peripheral blood infiltration is of prognostic value. In combination with immunocytochemistry, ultrastructural analysis can be a useful tool for the diagnosis of HCL variant.

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References

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