Enzyme histochemical analysis on cryostat sections of human bone marrow

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SUMMARY A recently developed procedure, that has been shown to be suitable for detailed immunohistological analysis, has been used to prepare cryostat sections of bone marrow to investigate whether enzyme-histochemical techniques are also feasible on such material. A selected group of enzymes, some of which are inhibited or destroyed in paraffin- or plastic-embedded samples, have been demonstrated. The morphological details obtained were satisfactory in the preparations. The enzymes were dipeptidyl(aminopeptidase IV (for T lymphocytes); tartrate-resistant acid phosphatase (for hairy cell leukaemia); acid phosphatase and non-specific esterase (for macrophages and monocytes); ATPase and S'nucleotidase (for B lymphocytes); and peroxidase or chloroacetate esterase (for granulocytic cells). In these preparations strong enzyme activities were shown. In adjacent sections the immunological analysis of membrane markers could also be performed contributing to a comprehensive study of the normal and malignant bone marrow cells.

Both histochemical1-3 and immunological methods4 are routinely used for the analysis of bone marrow aspirates in leukaemias and lymphomas. Similar studies performed on bone marrow biopsies could potentially be more informative because in histological preparations the tissue organisation is maintained. It is nevertheless very difficult to reconcile the different technical requirements which are necessary for optimal morphological, histochemical and immunological analysis of tissue biopsies. Some excellent enzyme histochemical studies have been already performed on bone marrow using cryostat and paraffin sections of prefixed, edetic acid decalcified tissue blocks,5-6 or methacrylate-embedded samples without decalcification.7,8 Nevertheless under these conditions sensitive enzymes such as magnesium-dependent adenosine triphosphatase (ATPase), S'nucleotidase (S'Nase) and dipeptidyl(aminopeptidase IV (DAP-IV) are inhibited or destroyed.9,10

Intracytoplasmic immunoglobulins can be successfully demonstrated on paraffin and plastic sections of bone marrow using immunoperoxidase technique.11 Nevertheless consensus has been recently reached about the fact that the staining of T and B lymphoid cells with antisera to membrane-associated antigens is severely handicapped by fixation and/or embedding procedures.12,13

Recently an interesting new development has taken place in the immunological analysis of malignant non-Hodgkin’s lymphomas. Some laboratories have emphasised that in properly processed frozen sections not only can excellent staining of membrane antigens be achieved but also the preservation of tissue morphology is acceptable.12,14,15 In line with this new trend we have demonstrated that bone marrow trephine biopsies embedded in gum-sucrose can be frozen and conveniently cut in a cryostat with the preservation of tissue (including membrane) antigenicity and morphology.16

In this paper we demonstrate that cryostat sections obtained with the same method (or using polyacrylamide gel as an alternative embedding medium) are also suitable for enzyme histochemistry. In addition to the enzymes which can be demonstrated in formalin-fixed paraffin- or plastic-embedded material (see Methods) a number of other enzymes could be detected in cryostat sections. These include Mg2+-dependent ATPase,17,18 S'nucleotidase 18 and dipeptidyl(aminopeptidase IV.10,19,20

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Material and methods

A few selected clinical cases have been studied as examples of the application of these methods.

PATIENTS

Three normal individuals (volunteers) and 12 patients were studied. The group of patients included eight cases of non-Hodgkin's lymphomas (NHL), one case of hairy cell leukaemia (HCL), two cases of B chronic lymphocytic leukaemia (B-CLL) and one case of T type CLL (T-CLL). In six out of the eight NHL cases the diagnosis had been made on lymph node histology (three cases of centrocytic-centroblastic lymphoma, one case of Burkitt-like lymphoma, one case of immunoblastic lymphoma and one case of unclassified lymphoma). In these six patients the trephine biopsy was performed as a staging procedure. In three no involvement was seen; in the other three (one case of centrocytic-centroblastic, one Burkitt-like and one unclassified lymphoma) the B lymphoid infiltration of bone marrow was identified. In the remaining two of the eight NHL cases the diagnosis of malignant lymphoma was first reached on the basis of the analysis of bone marrow trephine biopsy. One of these cases had a lymphoplasmocytoid lymphoma, diagnosed by classical histology in paraffin-embedded decalcified sections. This diagnosis was confirmed by the immunohistological analysis of frozen sections which demonstrated uniform λ+, μ+ populations. In the other case some lymphoid nodules were detected in the paraffin embedded biopsy but the definite diagnosis of malignant B cell proliferation was made by showing λ+ monoclonality of B cells on cryostat sections. In the three cases with CLL as well as in HCL the bone marrow involvement had been demonstrated by both histological and immunological analysis.

PREPARATION OF SAMPLES

Four cm long cores of bone marrow biopsies were obtained with an 11-gauge Jamshidi needle. Half of the sample was fixed in 10% formalin and embedded in paraffin after decalcification.21 The other half of the bone marrow core was submerged in gum-sucrose solution (1 g gum acacia + 30 g sucrose dissolved in 100 ml distilled water) as previously described10 or embedded in polyacrylamide gel (30% acrylamide, 1 ml; 1% bis-acrylamide, 1 ml; 0.5M Tris buffer pH 6-8, 2.5 ml; distilled water, 5:33 ml; 10% ammonium persulphate 0.05 ml; TEMED 0.005 ml) and snap-frozen in liquid nitrogen. These frozen cores of bone marrow samples were stored at −70°C until sectioning. In spite of the fact that no decalcification was performed no damage to the knife was evident when the sections (7–10 μm thickness) were cut in the cryostat. These sections were quickly dried on glass slides, fixed in neutral 10% calcium-formalin (4°C, 5 min), washed in distilled water and air dried with a fan. This preparation was used for all enzymes except DAP-IV where cold chloroform-acetone mixture was used as a fixative (5 min).

Reagents were from Sigma, St Louis, MO, with the exception of glycidyl-prolyl-4-methoxy-beta-naphthylamide purchased from BACHEM Feinchemikalien AG, Bubendorf/Schweiz.

The histochemical demonstration of the enzymes has been carried out as follows:

Non-specific esterase (NSE) with and without fluoride inhibition, using α-naphthyl acetate dissolved in ethylene glycol monomethylether (EGMME) (20 mg/0.5 ml), 0.067M neutral phosphate buffer (50 ml) and freshly prepared hexazonium pararosaniline (HPR) (2 ml) as a coupler. Incubation time 15–30 min, room temperature.22

Acid phosphatase (ACP) with and without L-tartrate inhibition, using naphthol AS-BI phosphate as substrate (10 mg dissolved in 0.5 ml EGMME), pH 5 acetate buffer (50 ml) and HPR (2 ml) as coupler; incubation time 1 hour at 37°C.23,24

Peroxidase (PX) using diaminobenzidine (20 mg dissolved in 0.5 ml EGMME),24 neutral Tris buffer (50 ml) and one drop 30% H2O2; incubation time 10 min, room temperature.25

Chloroacetate esterase (CAE), using naphthol AS-D chloroacetate (3 mg dissolved in 0.3 ml dimethylformamide) as substrate, neutral 0.067 M phosphate buffer (10 ml) and fast blue B (10 mg) as coupler; incubation time 15 min, room temperature.26

Alkaline phosphatase (ALP) using naphthol AS-BI phosphate (10 mg dissolved in 0.5 ml EGMME) as substrate, 0.1 M Tris buffer pH 9 (50 ml) and hexazonium new fuchsin (0.5 ml) as coupler; incubation time 30 min, room temperature.27

ATPase and 5′nucleotidase using modified lead-methods;14 incubation time, one hour, 37°C.

Dipeptidyl (amino)peptidase IV (DAP-IV), using glycidyl-prolyl-4-methoxy-beta-naphthylamide (2 mg dissolved in 0.2 ml dimethylformamide) as substrate, neutral 0.067 M phosphate buffer (5 ml) and fast blue B as coupler (5 mg); incubation time one hour, 37°C.10

After histochemical staining the sections were counterstained with haematoxylin when needed and mounted in glycerol-formalin (9:1).

Parallel sections have been fixed in ethanol and incubated for 30 min at 20°C with goat antibodies to human kappa and lambda light chain directly labelled with fluorescein isothiocyanate (FITC) and washed for a further 30 min prior to mounting in glycerol-formalin.
Fig. 1  Cryostat sections of normal bone marrow. In (a) and (b) non-specific esterase (NSE) staining is shown. In (b) the non-neoplastic lymphoid nodule is composed of a network of NSE positive cells (large arrow) amongst NSE negative or slightly positive lymphocytes. The processes of the NSE positive cells are similar to the network of follicular dendritic cells within the germinal centres of lymph node. Around the lymphoid nodule scattered strongly NSE reactive macrophages are also seen (small arrow). In the section adjacent to (b) the same lymphoid nodule has been stained for ATPase (c). The membranes of B lymphocytes are ATPase positive. No nuclear counterstaining was used. (a) × 250; (b) × 250; (c) × 400

Fig. 2  Cryostat sections of bone marrow with B-cell lymphoma (IgM+, lambda+ monoclonal population). (a) = haematoxylin and eosin stain (× 70); in (b) lymphoid cells on a serial section to (a) show ATPase positive membrane staining (× 250), no nuclear counterstaining; in (c) the same neoplastic nodule, on a serial section, immunostained for antilambda light chain (direct immunofluorescence method—FITC), photographed on a fluorescence microscope with filters for FITC. Virtually all B cells are stained. Similar staining was seen with anti IgM (μ specific) but staining with anti kappa light chain was negative (not shown). (× 400)

Results

PREPARATION OF BONE MARROW BIOPSIES

As previously reported gum sucrose embedding is important for obtaining good sections of frozen bone marrow biopsies. This is a thick fluid which can nevertheless easily penetrate into the bone marrow cavities and thus gives a compact homogeneous plasticity to the frozen samples. This is necessary for even cutting. The morphological quality of these sections is obviously inferior to those obtained from paraffin- or plastic-embedded tissues, but the details obtained in the haematoxylin-stained sections of the frozen material are surprisingly rich and adequate for certain diagnostic purposes (Figs 1 and 2). Using polyacrylamide gel as an alternative embedding medium thinner sections can be obtained improving the histochemical and immunohistological analysis.
ANALYSIS OF NORMAL AND NON-INVOLVED BONE MARROW SAMPLES

In these six specimens all enzymes studied were successfully demonstrated. Peroxidase (PX) and chloroacetate esterase (CAE) distinguished granulocytic cells from erythroid, megakaryocytic and monocytic cells which were mainly unreactive. Alkaline phosphatase (ALP) was positive in capillary endothelium and in fibroblast-like cells. Nonspecific esterase (NSE) was a suitable marker for megakaryocytes, monocytes and resident macrophages. These macrophages could be distinguished from monocytes by their irregular shape and more intense positivity (Fig 1). Fluoride abolished NSE reactivity in monocytes as well as in a large proportion of macrophages.

Lymphoid nodules were observed in two normal bone marrow samples. The lymphocytes in these nodules appeared to show slight reactivity for NSE. These lymphocytes formed contacts with processes of dendritic cells which had higher NSE activity. These dendritic cells had similar shapes and reactivity to the follicular dendritic cells seen in germinal centres of lymph nodes (Fig 1).

A somewhat similar histochemical pattern was observed with acid phosphatase (ACP) except that the expression of ACP activity was stronger in the disseminated bone marrow macrophages than in dendritic type cells or monocytes. L+tartrate completely inhibited the ACP reactivity in all of these cell-types except in a few macrophages and, when evident, some osteoclasts.

ATPase and 5' nucleotidase activities were mainly observed in the blood vessel walls and in the lymphocyte membranes within the lymphoid nodules (Fig 1c).

DAP-IV activity was confined to a few scattered T lymphocytes and to endothelial-like structures corresponding to the sinuses.

ANALYSIS OF MALIGNANT LYMPHOID INFILTRATION

The lymphoid cells had virtually no PX and CAE activity while the residual granulopoietic tissues were strongly positive for these enzymes. Thus the malignant infiltrations which obliterated the normal bone marrow structure could be clearly visualised (Figs 3 and 4).

Both the diffuse and nodular infiltrations were obvious. The NSE was unreactive or only slightly positive with lymphoma cells but a variable number of strongly NSE positive scattered “reactive” macrophages were seen amongst these neoplastic cells.

Certain enzymes positively identified the neoplastic lymphoid populations (Table). ATPase could be demonstrated on the malignant B lymphocytes in two cases of centroblastic-centrocytic lymphomas, one case of lymphoplasmocytoid lymphoma and in the two cases of B-CLL. This feature of the malignant B cells paralleled a similar ATPase reactivity on normal bone marrow nodules (Fig 1). The single case of T-CLL (and similarly normal T lymphocytes) were however ATPase-negative. In contrast DAP-IV was negative not only
The selectivity of enzyme activities demonstrated in biopsies of frozen bone marrow*

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*Alkaline phosphatase was demonstrated on blood vessels and fibroblast-like cells.
†DAP-IV activity was also present on sinus lining cells.
§Demonstrable as a “dot-like” positivity only on formalin fixed samples.
+/− Variable.

in residual normal bone marrow cells but also in all cases of B lymphoid malignancies. The one case of T-CLL studied was, however, strongly DAP-IV positive (Fig 5). 5′nucleotidase was negative or highly depressed in the neoplastic cells of all cases.

Finally PX, CAE, NSE, ATPase and 5′nucleotidase enzymes were all negative in the case of hairy cell leukaemia (HCL). Many HCL cells expressed ACP variable activity resistant to tartrate inhibition (TRAP) confirming the diagnostic specificity of this enzyme on HCL using cryostat sections (Fig 6).

Discussion

Our observations and the findings by other groups are briefly summarised in the Table. The enzymes chloroacetate esterase (CAE), non-specific esterase (NSE), peroxidase (PX) and both alkaline and acid phosphatase (ALP, ACP) can be demonstrated in fixed paraffin and methacrylate-embedded samples. Our results confirm previous observations about the distribution of these enzymes in the various normal and infiltrating cell types of the bone marrow. These enzymes together with the periodic acid-Schiff reaction are also used for the routine histochemical analysis of bone marrow smears.

The important point in the Table is that this range of “classical enzymes” as well as the more sensitive DAP-IV, ATPase and 5′nucleotidase can be detected in sections of frozen bone marrow biopsies with surprising ease and sensitivity. DAP-IV, ATPase and 5′nucleotidase are normally inhibited or destroyed by fixation and embedding procedures. Further details about the comparative sensitivity of histochemical methods applied on formalin-fixed paraffin-embedded and frozen tissue blocks have been described elsewhere.

Amongst the panel of reagents shown in the Table, the individual enzymes have distinct “roles” to play.
CAE and PX or ACP and NSE characterise granulopoietic cells and subpopulations of macrophages, respectively. These enzymes always clearly demonstrate the extent to which the bone marrow is replaced by neoplastic lymphoid cells. The other three enzymes, DAP-IV, ATPase and TRAP can positively identify whether these infiltrating cells are of T, B or hairy cell origin.

We feel that the relatively simple and economical methods shown here are valuable in routine histopathological analysis of bone marrow disorders for two main reasons. Firstly, as we have showed previously similarly processed bone marrow samples can be stained with various antisera for the detection of both membrane and cytoplasmic antigens. The uniform "monoclonal" light chain expression—that is, predominantly kappa or lambda, of the malignant B lymphoid population can be established. The new range of murine monoclonal antibodies for lymphoid subsets have also been applied with success in these adjacent sections. The histochemical methods are standard procedures and the monoclonal antibodies are becoming commercially available. It will, therefore, be possible, with the application of the described methods, to compare routinely the histochemical findings with the immunological phenotypes on normal and malignant cells. Secondly, parts of the bone marrow cores should be always preserved for routine histology. A further improvement is to cut the bone marrow cores longitudinally and process one part with plastic embedding (for optimal morphology) and the other part with gum sucrose embedding and freezing (for histochemistry and immunocytochemistry), as has been suggested by Bartl and his colleagues. In this way adjacent areas of optimally processed tissues can be analysed by complementary techniques.

Finally, we would like to call attention to the observation that 5′-nucleotidase is negative or weakly expressed in neoplastic B lymphocytes when compared to the strong expression of normal B cells in the follicle lymphocyte corona. This observation can already help to establish, together with the immunological analysis, whether the small focal nodules in the bone marrow of patients with lymphoma is aggregates of normal B cells or represent predominantly malignant populations. Nevertheless, the explanation of this phenomenon is unclear. It may represent malignancy associated changes in the expression of a membrane-linked enzyme, or may simply reflect that the 5′-nucleotidase negative B lymphomas derive from a B cell subset which is different from the corona B cell. As the analysis of subset heterogeneity (within normal B cell populations) by recently developed monoclonal antibodies has already been started it will soon be feasible to investigate this question further.

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