Morphology of lymphatic cells and of their derived tumours

FRANCO RILKE1, SILVANA PILOTTI1, ANTONINO CARBONE1, AND LUCIANO LOMBARDI2

From the 1Division of Anatomical Pathology and Cytology and 2Division of Experimental Oncology A, Istituto Nazionale per lo Studio e la Cura dei Tumori, Via Venezian 1, 20133 Milan, Italy

The morphological description of diseases of the lymphatic and reticulum cells, and the classification of neoplasms derived from these cells, are in a state of some confusion, which arises, in part at least, from the different approaches and terminology used by histopathologists and by cytologically orientated haematologists. The addition of electron microscopy, of enzyme histochemistry, and of immunological methods for cell typing has not entirely resolved the differences. The subject is still in a state of rapid development but an analysis of current knowledge is needed to promote mutual understanding among histopathologists, haematologists, immunologists, and many others who may be concerned. Whereas attempts to classify neoplastic disorders of the haematopoietic system have largely succeeded because the derivation of the tumour cells from the normal cells could be inferred from morphological study of marrow and blood cells, we have had to wait much longer for a comparable classification of lymphoreticular neoplasms—with the possible exception of Hodgkin’s disease.

In non-Hodgkin’s malignant lymphoma (ML) a major advance was made by Rappaport (1966; Rappaport et al., 1956), who moved from a popular terminology (lymphosarcoma, reticular cell sarcoma, and giant follicular lymphoma) to one based on lymph node structure, but the first step in the direction of a cytological analysis of the variety of ML was the use of haematological staining techniques applied to tissue sections in order to bridge the gap between histological findings and haematocytological data. For this purpose the use of the Giemsa stain was proposed by Lennert (1957, 1961). The major indications were, on the one hand, the morphological specificity of germinal centre cells and the comparability with them of the cells of follicular lymphomas and, on the other hand, the fact that follicular neoplasms made up of large and small cells were not tumours of different cell lines but of cells of the same line in different stages of modulation. Details of nuclear structure, nucleolar configuration, and the staining properties of the cytoplasm were also helpful in distinguishing ML with apparently similar morphology when stained with haematoxylin and eosin only. Further distinctions were derived from enzyme histochemistry applied to haematological and histological material (reviewed by Leder and Stutte, 1975; Harigaya, 1977), the significance of which was increased by the combination with electron microscopy (reviewed by Müller-Hermelink and Kaiserling, 1975; Kaiserling, 1977a).

However, the major recent contribution to the identification of the linkages between the cells of the normal lymphoreticular system and their malignant counterparts comes from immunological typing (reviewed by Preud’homme et al., 1975). The discovery of markers that are morphologically undetectable has permitted the characterisation and, to a great extent, also the function of both normal and malignant cells.

Since B and T lymphocytes (for reviews see Roitt et al., 1969; Henry and Goldman, 1975; Lennert and Müller-Hermelink, 1975) can be characterised by a variety of surface markers, the same techniques have been applied extensively to neoplasms of the lymphoreticular system (review, Stein, 1975). However, as has been stressed repeatedly (Brown et al., 1974; Seligmann et al., 1977a), great caution is advised both in the technical evaluation of the results and in the transfer of the data obtained from normal cells to pathological conditions. The remarkable progress in the immunological characterisation of ML is unfortunately occasionally hindered, particularly when detailed immunological analyses of valuable case material have been correlated with inadequate morphological investigations and inappropriate histological diagnoses.

The process of revision of the classification of ML

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has brought into focus in recent years the fact that these disorders are, in fact, neoplasms of the immune system and that the great majority of them are derived from lymphoid cells at various stages of functional and morphological modulation. More specifically, while some classification schemes stressed predominantly nondistinctive morphological features (Bennett et al., 1974; Dorfman, 1974), others linked morphological appearances to functional phenotype (Lukes and Collins, 1974). A cytogenealogical approach expressed, however, in purely morphological terms was preferred for the Kiel classification (Gerard-Marchant et al., 1974), which also incorporates a distinction between low- and high-grade ML. The existence of a group of ML with intermediate biological behaviour was demonstrated later on the basis of cell kinetic studies (Silvestrini et al., 1977). Finally, the classification supported by the World Health Organisation (Mathé et al., 1976) tried to express recent subdivisions while still respecting the conventional nomenclature.

Although a number of areas of uncertainty still remain, it seems more meaningful to support the natural trend towards the separation of nosographic entities. For this purpose, in the following review the strict adoption of any one classification is avoided, and within the limits of the data at present available the known entities are grouped according to the cell type of possible origin. The class of lymphocytic ML is heterogeneous, since the only common denominator is the small lymphocyte-like size of the cell, but this shortcoming is partly compensated by the existence of well-defined clinicopathological entities. For example, malignant lymphomas of immunoglobulin-secreting cells and of germinal centre cells represent excellent examples of the correlation that is possible between the normal cells and their malignant counterparts. Immunoblastic ML, could, to a certain extent, be considered together with ML of immunoglobulin-secreting cells; however, insufficient knowledge of T-type immunoblastic ML and the high percentage of large cell ML which are purely morphologically immunoblastic but 'receptor-silent', suggest a separate categorisation. The last group in which lymphoblastic ML and acute lymphoblastic leukaemia (ALL) have been combined depends much less on morphological and functional analysis than on immunological and biochemical properties. The comparatively exiguous group of neoplasms of histiocytic origin should not be classified among ML, since it would be preferable to reserve this term for malignancies of strictly lymphatic origin, except for Hodgkin's disease, which in turn deserves a fully autonomous position. The different cytogenealogy of reticulum cells in general, and of histiocytes in particular, makes this distinction necessary in spite of the close functional and anatomical relationship between the respective cell lines of origin.

This classificatory rearrangement (Stein, 1975) is based mainly on the work presented in recent years by Professor Lennert's group at the University of Kiel (Lennert et al., 1975b; 1975c) and has been terminologically revised (Gerard-Marchant et al., 1974). However, reference is made, whenever necessary, to other systems and especially to the Rappaport classification (Rappaport et al., 1956; Rappaport, 1966) because it has been widely accepted in many countries for many years by pathologists and clinicians.

The illustrations are from our own material, observed during the last four years, including over 500 cases of ML, 150 of which were also investigated ultrastructurally.

**Malignant lymphoma, lymphocytic**

**CHRONIC LYMPHOCYTIC LEUKAEMIA**

This disease, first described by Virchow (1864-65), is characterised by an excessive number of small lymphocytes in the blood and bone marrow, and in most patients also in lymph nodes, spleen, liver, and other organs (Wintrobe et al., 1974). Although its malignant nature is out of the question, its peculiar clinicopathological behaviour has been noted (Galton, 1966; Dameshek, 1967; Wintrobe et al., 1974), and clinical classification and staging systems take this into consideration (Levin et al., 1973; Rai et al., 1975; Binet et al., 1977a). In the majority of cases of chronic lymphocytic leukaemia (CLL) the proliferating cell is of B-cell origin (B-CLL), while in a small percentage it is of T-cell origin (T-CLL).

In sections of lymph nodes, B-CLL consists of a diffuse monomorphic proliferation of lymphocytes, most of which are morphologically not atypical and are almost indistinguishable from normal. The size of the latter is quite variable (Wintrobe et al., 1974; Kaung and Ott, 1975), but lymphocytes of CLL often give the impression of being slightly larger. Their nucleus is round with a regular contour and measures 7 to 9 μ in diameter; the chromatin is made up of coarse, mosaic-shaped chromocentres, and the nucleoli are either inconspicuous or hardly visible. Mitotic figures are rare. The cytoplasm is scanty, lightly basophilic, and PAS-negative, whereas in smears it occasionally reveals tiny granules and droplets of glycogen, which show a positive diastase-sensitive periodic acid-Schiff (PAS) reaction. The glycogen content is higher than in normal lymphocytes (Astaldi and Verga, 1957; Leder, 1971). The acid phosphatase reaction is weak to moderate in a minority of cells and less evident than in normal lymphocytes (Douglas et al., 1973; Catovsky et al.,
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1974a). Beta-glucuronidase is also scarce (Zittoun et al., 1973), whereas a strong membrane-bound ATPase reaction can be visualised (Kaiserling, 1977a). Both intracytoplasmic crystalline inclusions associated with IgM (Hurez et al., 1972; Clark et al., 1973), IgMk (Mennemeyer et al., 1974), and IgA (Cawley et al., 1973) and non-crystalline PAS-positive intracytoplasmic inclusions related to an excess of IgM in the neoplastic tissue (Stein et al., 1973) have been reported. It is questionable, however, whether these cases are true CLL or whether they should be considered lymphoplasmacytoid immunocytomas with a CLL-like clinical course.

While irregularly shaped lymphocytes of germinal centre origin (centrocytes) are essentially absent, in the tissue there is usually an admixture with a variable number of so-called prolymphocytes and lymphoblasts. The latter are up to twice the size of the small lymphocytes, and their nuclei contain sparse, finely granular chromatin, which is in part adherent to the nuclear membrane and in part arranged in threads that are extended between the nucleolus and the nuclear membrane. The large amphophilic nucleolus is central, and the cytoplasm shows a moderate degree of basophilia and contains no granules. These cells are also called paraimmuno blasts, because of their nuclear resemblance to immunoblasts, by those who reserve the conventional term 'lymphoblasts' for the immature lymphoid cell of ALL (Lennert, 1976). Prolymphocytes reveal morphological features that are intermediate between lymphocytes and lymphoblasts, and in sections both cell types are either irregularly scattered or more often grouped in pseudofollicular, clear 'proliferation centres' (Lennert, 1976) of variable size, which are not surrounded by condensed reticulin fibres as are the neoplastic follicles of follicular (nodular) ML. In these areas mitotic activity is increased. B-CLL lymphocytes replace the involved lymph nodes completely, and usually no remnants of normal structures are left; the capsule and the subcapsular sinuses are often still recognisable. Other cell types, such as plasma cells, lymphoplasmacytic cells, and follicular centre cells, are rarely found.

In the blood of the majority of patients with B-CLL, there are usually more than 10 × 10^9/l and less than 100 × 10^9/l lymphocytes at the time of diagnosis, with considerable fluctuations during the course of the disease. Another haematological definition of B-CLL is a persistent lymphocytosis of at least 5 × 10^9/l associated with a lymphocyte count of 25% or more in the bone marrow films (Galton, 1966). In blood smears the neoplastic, small lymphocytes dominate the picture, and there is no morphological abnormality to distinguish them from normal lymphocytes. The nuclei contain dense chromatin which is subdivided into coarse blocks, nucleoli are therefore unidentifiable. Nuclear abnormalities are uncommon in B-CLL, but the cells are fragile, as is apparent from the large number of damaged cells in the smears (Galton, 1966). The narrow cytoplasmic rims are pale blue and usually devoid of granules and vacuoles. In a number of otherwise typical cases of CLL, large (up to 20 μ in diameter) lymphocytes may predominate, and their prognostic significance is a matter of debate (Gray et al., 1974; Peterson et al., 1975; Binet et al., 1977b).

Immature cells (lymphoblasts) are more commonly detected in lymph node aspirates and sections than they are in blood or marrow films or marrow sections prepared at the same time (Galton, 1966). Marrow invasion is always an early event (Galton, 1966), as revealed by aspirate smears; however, marrow biopsies show variable pictures of focal, nodular, diffuse, or massive invasion (Duhamel, 1974), which correlate to a certain extent with the indolent or active clinical behaviour of the disease (Gray et al., 1974; Carbone et al., 1978).

A minority (1-3%) of patients with CLL may develop an acute leukaemia after a variable, but usually long (more than 5 years), interval (Zarrabi et al., 1977). Either truly undifferentiated cells or large lymphoblasts with vacuolated cytoplasm containing rare granules are found to circulate in the blood and to replace to a great extent the marrow. The finding of surface immunoglobulin on the blasts of two cases, and in one of them of the same monoclonal IgM with an anti-IgG antibody activity on both the leukaemic lymphocytes and the blasts cells, is against the possibility of a second malignancy and is suggestive of a derivation of the blasts from the same clone as the small lymphocytes (Brouet et al., 1973b).

In lymph node sections the numerical increase of lymphoblasts among the monomorphic B-CLL lymphocytic population is usually considered by histopathologists to be suggestive of neoplastic progression. The development of an anaplastic lymphoid malignancy ('reticular cell sarcoma') during the terminal phases of CLL was described by Richter in 1928. It has distinct clinicopathological features (Long and Aisenberg, 1975) and is usually diagnosed at necropsy. Rather than a metachronous association of two malignant lymphomas, it should be considered an immunoblastic sarcomatous evolution of CLL with predominant tissue manifestation. The possible role of long-term chemotherapy in the switch-on of the blastic phase is at present a matter of debate. By contrast, the development of acute megaloblastic or myelomonocytic leukaemia is an exceedingly rare event (cf. myeloma).

Ultrastructurally, the great majority of the
lymphocytes (Figs 1 and 2) have round nuclei with coarsely clumped chromat in (Mori and Lennert, 1969; Cawley and Hayhoe, 1973). There are no indentations of the nuclear membrane; however, occasional nuclear pockets are found. The inconspicuous nucleoli frequently have a characteristic ring-shaped pattern. Conversely, the lymphoblasts (Fig. 3) have diffuse nuclear chromat and well-developed nucleoli with a prominent nucleolomema. The prolymphocytes (Fig. 2) have a roughly oval nucleus and a well-developed nucleolus as in the homonymous cells of Galton's prolymphocytic leukaemia (PL) (Galton et al., 1974), but they have a more abundant cytoplasm and a less clumped nuclear chromatin. Prolymphocytes of both B-CLL and PL are distinguishable from centrocytes, which have deeply indented irregular nuclei, inconspicuous nucleoli, and scanty cytoplasm with scarce organelles.

The cytoplasm of B-CLL cells is rather scanty in lymphocytes and is somewhat more conspicuous in lymphoblasts and prolymphocytes. It contains a small or moderate number of mitochondria, numerous free ribosomes, a small Golgi apparatus, and a few rough endoplasmic reticulum cisternae. Abundant cytoplasmic microfilaments are present in some cells. The previously mentioned inclusions with a regular crystalline-like structure lie within the rough endoplasmic reticulum cisternae. A morphological marker that indicates a plasmacytoid transformation of rare CLL lymphocytes is represented by some cisternae of long, flat, and rough, endoplasmic reticulum placed around the nucleus (Fig. 4). Scanning electron microscopy (SEM) does not permit the identification of the lymphocytes of B-CLL as neoplastic (Braylan et al., 1976).

Extensive studies of cell surface immunological markers, such as surface-bound immunoglobulin (Pernis et al., 1970), membrane receptors for the activated third component of complement (Bianco et al., 1970; Ross et al., 1973), and aggregated IgG (Dickler et al., 1973) on CLL lymphocytes (Johansson and Klein, 1970; Grey et al., 1971; Wilson and Nossal, 1971; Aisenberg and Bloch, 1972; Frölund et al., 1972; Preud'homme and Seligmann, 1972; Piessens et al., 1973; Ross et al., 1973) proved that this B-lymphocyte proliferation is made up of cells in most of which no immunoglobulin secretion takes place but which do bear synthesised membrane-bound immunoglobulin. This is of one type only and is usually restricted to one type of heavy chain (μ, γ, or α) and one light chain (κ or λ), thus indicating the monoclonality of the cell proliferation (Aisenberg and Bloch, 1972), even though biclonal processes have rarely been detected (Preud’homme and Seligmann, 1972). In a number of cases δ chains are associated with μ chains, as may be observed in normal lymphocytes (Fu et al., 1974). Routine immunofluorescent staining procedures for the demonstration of immunoglobulin on the cell surface of leukaemic lymphocytes reveal positivity that is weak (Braylan et al., 1976), particularly when compared with that of the cells of lymphosarcoma-cell leukaemia (Aisenberg and Bloch, 1972; Aisenberg et al., 1973a), that is, the leukaemic counterpart of a lymphoma of germinal centre cell origin. In general, the cells of B-CLL also have weakly represented complement receptor sites (Shevach et al., 1972) and membrane receptors for aggregated IgG (Braylan et al., 1976). However, the expression of binding sites varies considerably from case to case. The absence of surface immunoglobulin (Piessens et al., 1973; Wilson and Hurdle, 1973) has been related to the proliferation of a pre-Ig-synthesising lymphocytic population (Preud'homme and Seligmann, 1972). B-CLL cells either fail to respond or respond poorly to mitogens in short-term cultures (Quaglin and Cowling, 1964). It has also been shown that a high percentage of them form rosettes with mouse erythrocytes (Catovsky et al., 1976; Koziner et al., 1977) in contrast to the cells of lymphosarcoma-cell leukaemia.

It has been suggested that B-CLL most likely represents the proliferative disease of a ‘virgin’ (B1) B-lymphocyte clone, which is still untouched by the antigen, is blocked in its modulation process, and is therefore unable to secrete immunoglobulin; but its derivation from ‘memory’ (B2) B-lymphocytes in some cases cannot be ruled out (Salmon and Seligmann, 1974). It has also been suggested that B1-cell proliferations are those whose cells bear surface immunoglobulin and have complement receptors, whereas B2-cell proliferations also bear surface immunoglobulin but are devoid of complement receptors (Stein, 1975).

When the tissue manifestation of B-CLL is not accompanied by an excess of cells in the blood and/or marrow, the disease has been referred to as well-differentiated, lymphocytic, diffuse ML (Pangalis et al., 1977); a leukaemic manifestation may appear later (Galton, 1966) or never (Goldberg and Emanuel, 1964). However, the morphology and the surface characteristics of the cells are the same as they are in B-CLL (Huber et al., 1974; Peter et al., 1974; Aisenberg and Long, 1975; Brouet et al., 1975b; Braylan et al., 1976). One major difference between B-CLL and well-differentiated lymphocytic diffuse ML was reported to consist of a less evident degree of hypogammaglobulinaemia in the latter (Pangalis et al., 1977). In spite of clinical and haematological differences, leukaemic and non-leukaemic B-lymphocytic neoplastic proliferations are so
Fig. 1 Chronic lymphocytic leukaemia. Small lymphocytes and a lymphoblast (L). (× 4000)

Fig. 2 Chronic lymphocytic leukaemia. A small lymphocyte (A) with coarsely clumped nuclear chromatin, numerous monoribosomes, scarce rough endoplasmic reticulum, and lysosome-like dense granules. A prolymphocyte (B) with a well-developed nucleolus and a larger cytoplasm. (× 10 000)
Fig. 3  Chronic lymphocytic leukaemia. A lymphoblast with diffuse nuclear chromatin and a prominent nucleolus. Some long rough endoplasmic reticulum cisternae and mitochondria are present. (× 9000)

Fig. 4  Chronic lymphocytic leukaemia. A small lymphocyte with a well-developed rough endoplasmic reticulum. (× 11 000)

Fig. 5  Malignant lymphoma, T-zone type. A T-associated plasma cell with a prominent nucleolus and a well-developed rough endoplasmic reticulum. (× 28 000)
closely related that they are probably just different haematopathological expressions of the same disease.

In large series of cases of CLL, occasional patients may be encountered in whom morphological and/or immunological deviations from the classic picture are noticed. Morphological deviations are represented by cases in which circulating cells have either cleft nuclei similar to Rieder cells and to small follicular centre cells (centroytes) of lymphoplasmacytic features (Rudders, 1976) and contain cytoplasmic inclusions. In general, lymphocytes of abnormal appearance have been related to a more rapid course of the disease. Along these lines, immunological variations are exemplified either by cases whose cells demonstrate a surface marker phenotype, which is comparable to that usually displayed by follicular centre cells (Braylan et al., 1976) or by so-called 'atypical CLL' cases whose cells contain intracytoplasmic IgG (Rudders, 1976). On the basis of these descriptions one might suspect that these atypical cases of CLL are, in fact, leukaemic manifestations of ML, respectively of follicular centre cell origin or of lymphoplasmacytoid type. In rare instances, both B- and T-cell surface markers have been found on leukaemic cells (Chin et al., 1973); however, their true B + T nature has been questioned (Brouet et al., 1975c; Siegal et al., 1976). Shifts of markers have also been noticed during the course of the disease (Kay et al., 1974).

T-CLL (Bentwich and Kunkel, 1973; Dickler et al., 1973; Lille et al., 1973; Wilson and Hurdle, 1973) is a rare entity (less than 2% of all CLL), with onset in adulthood, frequent massive splenomegaly, skin lesions, and variable lymphocyte count, which is characterised cytologically by somewhat larger lymphocytes than those of B-CLL (Brouet et al., 1975a). The cells have fairly large, occasionally deeply basophilic, PAS-negative cytoplasm which contains large azurophilic granules and shows a positive beta-glucuronidase and acid-phosphatase reaction (Catovsky, 1975). The nuclei contain coarse chromatin granules and a small nucleolus. Prolymphocytes and lymphoblasts are absent.

On electron microscopy the nuclei are irregular and there is a high content of lysosomal enzymes. Transmission and SEM cannot detect any morphological difference either between normal B and T lymphocytes (Alexander et al., 1976; Newell et al., 1976; Polliack, 1977) or between CLL with T or B surface immunological markers. Immunologically, the presence of receptors for sheep erythrocytes (Lay et al., 1971; Jondal et al., 1972) and the reactivity with heterologous anti-T-cell antiserum (Aisenberg et al., 1973b) indicate the T-derived nature of the cells.

Additional cases of this or of a closely related disease with a poor prognosis have been reported in Japan. The circulating lymphocytes, however, showed a higher degree of nuclear pleomorphism and fewer cytoplasmic granules (Uchiyama et al., 1977) as compared with the T-CLL described in western countries. As far as we are aware, detailed histological descriptions of lymph nodes involved by T-CLL are lacking.

T-ZONE MALIGNANT LYMPHOMA
Whether T-CLL bears any relationship to the T-zone lymphocytic ML has not yet been established (Lennert et al., 1975b). It seems, however, that this recently described entity (Lennert et al., 1975b; Lennert, 1976) is a well-defined one and that it represents neoplastic malignant proliferation of the lymphocytes of the nodal paracortical area. In fact, in cases that are not too far advanced, uninvolved secondary follicles which are widely dissociated by the neoplastic growth remain recognisable. The neoplastic cell population is somewhat pleomorphic and is made up of irregularly shaped lymphocytes, the nuclei of which are on average less hyperchromatic than are those of B-CLL. The chromocentres are smaller and not clumped, and a small prominent nucleolus is usually visible. The mitotic index is low. The cells display a strong acid phosphatase activity (Kaiserling, 1977b). The histological picture is also characterised by the presence of the normal components of the T-dependent paracortical area (Kaiserling, 1977b), that is, postcapillary venules lined by hob-nailed endothelial cells and surrounded by small lymphocytes, a network of reticulin fibres, and interdigitating reticulum cells, the clear, highly irregular nuclei of which may be recognised even on light microscopy (Veldman, 1970). These cells are considered to be the characteristic reticulum cells of the paracortical area of the lymph nodes. They do not usually display phagocytic activity. In addition, so-called T-associated plasma cells are part of the picture (Lennert et al., 1975a).

The intermingled rare large blast cells are similar to Hodgkin's mononuclear cells and may possibly be interpreted as T immunoblasts. Multinucleate cells of the same type simulate Reed-Sternberg cells. Six cases of a very similar entity, defined as ML of peripheral T-lymphocyte origin, were recently reported by Waldron et al. (1977) in elderly patients. The majority of the malignant cells of these cases were identified immunologically as T lymphocytes.

On electron microscopy the cellular population of the T-zone ML diffusely infiltrates the lymph nodes and consists of pleomorphic lymphoid cells, a smaller number of non-neoplastic macrophages, rare plasma cells, and granulocytes (Waldron et al.,
formed lymphocytes whereas cytotes, abundant cytoplasm and scarce rough endoplasmic reticulum cisternae, a variable number of mitochondria, and occasional lysosome-like dense granules located in the Golgi area. A few abnormal cells (Fig. 7) may show either a huge central nucleolus with a very prominent nucleolonema or an irregularly indented nucleus (Fig. 8) with dispersed chromatin; these cells recall the descriptions of mononuclear Hodgkin's cells (Dorfman et al., 1973; Glick et al., 1976). The identification within the tumour tissue of interdigitating reticulum cells can be considered a diagnostic marker of the T-zone lymphoma and other T-derived lymphomas (Kaiserling, 1977b). Interdigitating reticulum cells have a clear nucleus with scarce heterochromatin placed along the nuclear membrane and with a regular shape or, occasionally, with deep indentations. A small nucleolus is observed. The abundant cytoplasm is extended into interdigitating processes. No junctional differentiation is evident among the plasma membranes of two contiguous reticulum cells. In the cytoplasm a complex of vesicles and sacs, markedly contrasted by silver-methenamine staining, is found. This electron microscope cytochemical method is considered specific for glycoproteins (Veldman, 1970). The Golgi apparatus also shows a positive silver-methenamine reaction, which supports a functional correlation between the two structures. A few rough endoplasmic reticulum cisternae, some mitochondria, bundles of filaments, and a small number of monol- and polyribosomes are also found. No digestive vacuoles are seen.

PROLYMPHOCTYIC LEUKAEMIA (PL)

This disease (Galton et al., 1974) is a rare variant of CLL with distinctive features; lymphadenopathy is inconspicuous, whereas the spleen and liver are markedly enlarged. There is usually a high WBC count, and the proliferating cell has been designated 'prolymphocyte', with a different meaning, however, from that used in the WHO classification of neoplastic lymphoid diseases. The leukemic cells are larger than the lymphocytes of B-CLL, have a fairly large basophilic cytoplasm, a nucleus with condensed chromatin, and a single vesicular and prominent nucleolus. A variable number of lymphocytes as well as larger blast-like cells are also present. The latter are different from the lymphoblasts of CLL because of the larger size and the greater amount of chromatin. Abnormal looking cells are often found. Most of the reported cases have been of B-cell origin, but a few have been of T-cell origin. The former reveal a stronger expression of surface membrane-bound immunoglobulin and of complement receptors than the lymphocytes of B-CLL (Catovsky et al., 1973), while the T-cell prolymphocytes lack those markers but yield a high spontaneous E-rosette count. In smears, a high percentage of the cells of B-type PL contain glycogen (Stathopoulos et al., 1974) in the form of granules which are coarser than those found in B-CLL lymphocytes. Acid phosphatase content is weak but occasionally tartrate-resistant (Catovsky et al., 1974a). On the other hand, T-cell type PL cells reveal a higher content of acid phosphatase and are PAS negative.

Essential differences between the ultrastructural features of prolymphocytes and those of B-CLL lymphocytes consist of the larger size and the more prominent nucleolus, with the well-developed nucleolonema of the former. Ring-shaped nucleoli are seldom seen. In the cytoplasm a variable number of mitochondria and some rough endoplasmic reticulum cisternae are found. The blasts of PL have large nucleoli and dispersed chromatin. In comparison to prolymphocytes of B-CLL, prolymphocytes of PL are characterised by a clumped chromatin pattern and may therefore be placed morphologically in an intermediate position between small lymphocytes and the intermediate cells of CLL. No morphological differences have been reported between T- and B-type PL.

MYCOSIS FUNGIOIDES AND SÉZARY SYNDROME

Mycosis fungoides is an uncommon (less than 1% of all ML) but distinctive histopathological and clinical entity which involves primarily the skin (Edelson et al., 1974; Cline, 1975) and later, in over two-thirds of cases, the lymph nodes and other organs (Long and Mihm, 1974; Rappaport and Thomas, 1974). The cellular composition of the infiltrates of mycosis fungoides is lymphocytic in nature (Crosen et al., 1971); the cells are unique and different from those of other ML, and are considered to originate from T-dependent lymphocytes.

Histological diagnosis is usually made on skin biopsies of the second (indurated neoplastic cutaneous plaque) and of the third (myotic or neoplastic) stage of the disease. The neoplastic cellular population is identical in the cutaneous and extracutaneous sites. In the skin there is an infiltration by mononuclear atypical cells, which invade the papillary dermis and the basal layers of the
Fig. 6  Malignant lymphoma, T-zone type. Large lymphoid cell with two prominent nucleoli, many polyriboosomes, scarce rough endoplasmic reticulum, a small lipid droplet (L), and a few lysosome-like dense bodies. (× 8000)

Fig. 7  Malignant lymphoma, T-zone type. A tumour cell with a roughly oval, pale nucleus and a very large central nucleolus. (× 14 000)

Fig. 8  Malignant lymphoma, T-zone type. A neoplastic cell with an irregular, indented nucleus, abundant polyriboosomes, and small lysosome-like dense bodies. (× 8000)

Fig. 9  Mycosis fungoides: imprint of skin lesion. Medium-sized neoplastic cells with highly irregular and cerebriform nuclei (arrow). The nucleolus is hardly visible and the cytoplasm is clear. (Papanicolaou × 1000)
epidermis, in which the so-called Darier-Pautrier's abscesses are often formed within epidermolytic spaces. The neoplastic infiltrate consists of cells with a wide range of sizes. The most common cells, however, are medium-sized and measure 10 to 20 μ in diameter; they have a scanty, faintly stained cytoplasm, which shows a focal acid phosphatase activity (Schwarze, 1975) and contains perinuclear diastase-resistant, PAS-positive granules. Their nuclei are pleomorphic, with dense chromocentric chromatin and a small nucleolus.

The atypical cells may show a certain variation in size and shape with marked atypicalities. The larger the cells the more evident are the irregularities of the nuclear membrane, particularly in imprints (Fig. 9), with hyperconvolution and cerebriform patterns. In the very small cells nuclear details are barely visible. The so-called mycosis cells are a rare finding; they are the largest of the whole neoplastic cell population and show marked hyperconvolution and hyperchromasia of the nucleus and amphiphilic cytoplasm. In addition, a large number of abnormal looking histiocytes may be present (Robinowitz et al., 1976).

In lymph nodes, the neoplastic proliferation of mycosis fungoides seems to involve initially the 'thymic-dependent' paracortical area and later, to a variable extent, the remaining node (Thomas and Rappaport, 1975). All cell types present in the cutaneous infiltrate can be encountered in extracutaneous sites of invasion. In the spleen, either scattered foci in both the red and the white pulp or a diffuse infiltration may be seen (Variakojis et al., 1974). The selective involvement of the thymic-dependent periarteriolar lymphatic sheath has been stressed (Thomas and Rappaport, 1975).

Sézary's syndrome (Sézary and Bouvrain, 1938), a chronic leukaemia associated with erythroderma, is considered the leukaemic variant of mycosis fungoides (Lutzner et al., 1975; Robinowitz et al., 1976). In fact, the cutaneous infiltrate in Sézary syndrome is very similar to that of mycosis fungoides, and, on the other hand, in a high percentage of cases of mycosis fungoides circulating Sézary cells were found (Moran et al., 1977). In addition, in Sézary syndrome the earliest infiltration in lymph nodes was reported to be found histologically in tertiary follicles (Lennert, 1974). It was therefore proposed that mycosis fungoides and Sézary syndrome should be grouped together under the term 'cutaneous T-cell lymphoma' (Schein et al., 1976).

The circulating cells, which represent the diagnostic marker of the disease—even though not an absolute one (Lutzner et al., 1975)—are variable in number (between 10 and 50% of the nucleated cells in blood smears) and reveal nuclear and cytoplasmic features that are closely related to those described for mycosis cells. In marrow smears Sézary cells are rarer than expected in a leukaemic disease. The cells usually measure between 15 and 20 μ in diameter and have a high nuclear:cytoplasmic ratio. The scanty cytoplasm is well defined, appears light blue with the Giemsa stain, and does not contain granules, while empty vacuoles may be present. In thin blood smears the nuclei disclose their highly characteristic structure: the nuclear membrane shows various furrows, indentations, folds, and convolutions. Nucleoli are barely visible. The same finding is revealed by imprints of skin lesions as well as of lymph nodes that are involved by mycosis fungoides and by Sézary syndrome. Sézary syndrome also has a more markedly leukaemic small cell variant that is made up of lymphocytes measuring about 8 μ in diameter which are morphologically more closely similar to small lymphocytes even though nuclear indentations are present (Lutzner et al., 1973). The large cells have near-tetraploid DNA values and near-tetraploid chromosome counts, whereas the small cells have diploid DNA values and pseudodiploid or hyperdiploid chromosome counts.

Sézary cells reveal a granular acid phosphatase content, are moderately positive for β-glucuronidase and α-naphthyl acetate esterase, and are negative for peroxidase, naphthol-AS-D-chloroacetate esterase, alkaline phosphatase, adenosine-triphosphatase, Oil-Red O, and Sudan black-B (Löffler, 1972). PAS stains cytoplasmic granules of neutral mucopolysaccharides in a variable number without diffuse background staining (Crossen et al., 1971). Large and small Sézary cells have T-cell markers; they form spontaneous E rosettes, react with antihuman T-cell antiserum, respond to phytohaemagglutinin (Brouet et al., 1973a; Lutzner et al., 1973), and lack B-cell markers; they do not bear complement receptor sites and are devoid of surface-bound immunoglobulin (Broome et al., 1973). In addition, evidence was given that Sézary cells represent a neoplastic proliferation of T-helper cells (Broder et al., 1976). A few cases of Sézary syndrome in which the neoplastic cells failed to form E rosettes have also been reported (Goldstone et al., 1976).

Ultrastructurally, tissue-bound mycosis fungoides cells (Lutzner et al., 1971) are also very similar to circulating Sézary cells (Lutzner and Jordan, 1968; Zucker-Franklin et al., 1974). The nuclear:cytoplasmic ratio is high, and the nucleus is characteristically irregular, serpentine, indented, and lobed. However, this degree of irregularity varies largely from cell to cell and from case to case (Rosas-Uribe et al., 1974). Heterochromatin is abundant and concentrated along the nuclear membrane. One or two nucleoli with nucleolonema or ring-shaped
nucleoli and occasional nuclear inclusions are observed. The cytoplasm contains glycogen and a prominent network of microfilaments (Zucker-Franklin et al., 1974). The number of mitochondria, multivesicular bodies, rough endoplasmic reticulum cisternae, and free mono- and polyribosomes varies from case to case. Cytoplasmic pseudopods and lysosomes are found, whereas phagocytosis is never seen.

At SEM most of the Sézary cells show a moderate to markedly villous surface and do not display ruffled membranes (Polliack et al., 1977). Since Sézary-like cells are found in patients with a variety of nonlymphomatous dermatoses, it is of diagnostic importance that atypical cells should be demonstrated in clusters or sheets in the affected tissues.

**HAIRY CELL LEUKAEMIA**

This uncommon form of a chronic leukaemic lymphoproliferative disorder is characterised clinically in most cases by splenomegaly, hepatomegaly, and pancytopenia. Lymphadenopathy is a rare and late symptom. In sections of spleens there is a diffuse proliferation of monomorphic cells which measure between 12 and 20 μ in diameter. The nucleus is roundish or oval, occasionally kidney-shaped, often indented and wrinkled, and eccentric. It occupies about half of the cell area. Mitotic figures are rarely seen. The chromatin is delicate and loose, and nucleoli (if discernible) are small and pale. The cytoplasm is quite large, ill-defined, irregular, and lightly basophilic (grey-blue in Wright-Giemsa smears) and does not contain granules. Large blast-like cells are never found. In ordinary blood smears in which the cell morphology is best observed, the hairy finger-like projections of the cytoplasm may be recognised. However, they appear to be better preserved and defined in phase-contrast microscopy. In thin, well-stained smears and splenic imprints, small, cigar-shaped or roundish pyroninophilic cytoplasmic inclusions may be detected that correspond to the ribosome-lamellar complexes (Katayama et al., 1973).

The cytological marker is the presence of a prominent focal acid-phosphatase activity, which is resistant to degradation by tartaric acid (isoenzyme 5) (Yam et al., 1972). Even though hairy cells may occasionally be tartaric-acid sensitive, this isozyme should always be sought, preferably in blood smears (Katayama and Schneider, 1977); it is absent in normal lymphocytes but may be found in some CLL and PL lymphocytes, and its appearance in hairy cells should be considered as a newly acquired property that is related to the malignant status of the cells. Hairy cells are negative for Sudan black-B, peroxidase, and PAS, even though in some cells a PAS-positive granulation has occasionally been seen (Haak et al., 1974). A moderate granular sodium fluoride-resistant naphthol AS-D-acetate esterase activity, allegedly stronger than that of lymphocytes and weaker than that of histiocytes, has been reported (Flandrin et al., 1973). Recent extensive enzymohistochemical investigations revealed that hairy cells show only occasionally a-naphthyl acetate esterase activity, while they are negative for alkaline phosphatase, a-naphthylbutyrate esterase, sodium fluoride-resistant naphthol AS-D-acetate esterase, S'-nucleotidase, and N-acetyl-β-glucosaminidase (Nanba et al., 1977b).

The earliest and most striking proliferation of cells takes place in the spleen, which usually becomes markedly enlarged. Grossly, the spleen is dark red in colour, firmer than normal, and does not reveal discrete tumour masses. In the early stages, however, hairy cells appear first in the cords of the red pulp, and subsequently the sinusoidal structure and the Malpighian corpuscles of the white pulp tend to disappear because of compression and atrophy. In addition, an increase of actively phagocytosing histiocytes has been described around the arteries of the red pulp (Nanba et al., 1977b). Plasma cells, both normal and of atypical appearance, are found among the hairy cells. Their purely reactive nature has been questioned. Distended spaces filled with erythrocytes and lined by hairy cells and not by endothelial cells, which show a strong fluoride-sensitive naphthol-AS-D-acetate esterase activity, were recently described as pseudosinuses (Nanba et al., 1977c).

Marrow invasion may be either massive or focal. The marked and diffuse increase in reticulin fibres (Burke et al., 1974) explains the ‘dry tap’ that is most often found; however, imprints of marrow biopsies yield adequate diagnostic cytological material (Krause et al., 1977). In the liver, the invasion usually follows the sinusoidal structure, and pseudoangiomatosus lesions similar to those of the spleen and lined by hairy cells have also been described. In the lymph nodes, the neoplastic proliferation appears late in the course of the disease and begins in the B-cell region of the outer cortex with early infiltration of the marginal sinus. The spread to the paracortical and medullary areas develops later (Lennert, 1974). Lymph node invasion is often only partial, with residual follicles and foci of lymphoid cells.

The large number of synonyms used so far for the definition of this disease, which is still also called leukaemic reticuloendotheliosis (Ewald, 1923; Bouroncle et al., 1958), indicates the uncertainties about the proliferating cell type, the normal counterpart of which has not yet been identified. It has been questioned whether hairy cell leukaemia is really a
completely homogeneous disease rather than a group of subentities (Golde et al., 1977b), and its origin from histiocytic, endothelial, or lymphoid cells was and is still a matter of controversy.

The moderate nonspecific esterase activity, the occasional and inconsistent ability of phagocytic activity in vitro (latex particles and complement-coated zymogen particles) (Boldt et al., 1977) and in vivo (Catovsky et al., 1975a; Nanba et al., 1977b; Utsinger et al., 1977), and adhesion of the cells to glass and nylon (Flandrin et al., 1973; Boldt et al., 1977) are weak arguments, indicating that hairy cells may be monocytic in nature; and the severe monocytopenia associated with the disease has been interpreted in the same way (Seshadri et al., 1976). However, none of these is a compelling reason and furthermore, hairy cells do not produce muramidase (Catovsky et al., 1975a). An origin from endothelial cells also seems unlikely because of the tartric acid-resistant acid phosphatase activity and the absence of naphthol AS-D-acetate esterase reaction.

On the other hand, arguments identifying the hairy cell as a neoplastic B-modified lymphocyte, in addition to the morphological findings are: frequent surface-bound ATPase activity (as in CLL) and β-glucuronidase activity (as in malignant plasma cells) (Nanba et al., 1977b); the finding of surface immunoglobulin (Catovsky et al., 1974c; Stein and Kaiserling, 1974; Leech et al., 1975b; Boldt et al., 1977), probably IgM (Debusscher et al., 1975), and of intracellular immunoglobulin (Leech et al., 1975b; Golde et al., 1977a), probably IgMA (Debusscher et al., 1975); and reactivity with the Merritt B-cell alloantibodies (Naeim et al., 1977). However, surface immunoglobulin was demonstrated occasionally to be polyclonal on hairy cells, and receptors for cytophilic antibodies were identified (Jaffe et al., 1974b), while the search for complement receptors gave predominantly negative results (Burns et al., 1977). The responsiveness to mitogens was found to be impaired but less than it is in CLL (Haak et al., 1974). On the hairy cells surface immunoglobulin redistribution was induced by antibodies; polar cap formation occurs as with other B cells and was found to be very active and resistant to various agents, such as low temperature and sodium azide (Salsano et al., 1978). No evidence for a T-cell origin has been found so far.

Ultrastructural investigations also confirm that in lymph nodes the neoplastic cell proliferation begins in the outer cortex. The tumour population consists of one cell type only: the hairy cell shows the same features in all sites and is characterised by the abundant hairy-like cytoplasmic projections interdigitating in a complex way and giving to the cellular clusters a syncytial appearance. Two types of cytoplasmic projections are present on the cell surface (Katayama and Schneider, 1977): thin finger-like microvilli (Fig. 10) and large pseudopods (Fig. 11). Both projection types are present on the same cell, but the numerical ratio between microvilli and pseudopods changes from cell to cell.

The nucleus is polymorphous, either oval or reniform (Fig. 11), with slight indentations, or more often bilobed or multilobed. Nuclear pockets are rarely seen. Chromatin is more dispersed than it is in lymphocytes, and one or rarely two nucleoli are present. The ribosome-lamellar complexes (RLC) already mentioned are present in the cytoplasm of hairy cells in a variable percentage of cases. In different patients the RLC are present in 0-2% to 90% of the tumour cells (Katayama and Schneider, 1977). The RLC (Fig. 10) are 8 to 13 μ long and 0-5 μ wide, open tubular structures with a wall formed by concentric lamellae and ribosomes. RLC are rarely observed in other haematopoietic disorders, such as CLL, multiple myeloma, Waldenström’s disease, acute monoblastic leukaemia, centroblastic-centrocytic ML, and also in human adrenal cortex adenoma cells, monkey renal tubular cells, and plant cells (Anday et al., 1973; Katayama et al., 1973; Burke et al., 1976; Reynes and Diebold, 1977). However, since the RLC are much more numerous in hairy cells, they can be considered a useful diagnostic marker for light and electron microscopy.

Other features in the cytoplasm of hairy cells (Figs 10 and 11) are a limited number of mitochondria, a relatively well-developed Golgi apparatus, scattered rough endoplasmic reticulum cisternae, numerous free mono- and polyribosomes, pinocytotic vesicles, azurophilic granules, rare lysosomal inclusions, bundles of microfilaments, microtubules, centrioles, and occasional lipid inclusions. By SEM, hairy cells demonstrate microvilli, which are a peculiar characteristic of monocytes. A surface variation is manifested. Some cells have a villous surface whereas others have ruffled membranes only (Deegan et al., 1976); hybrid cells showing both features are also seen.

In summary, hairy cells show hybrid features that are predominantly typical for B lymphocytes and, in part, for monohistiocytic cells. A partial explanation may be that hairy cell leukaemia is not a completely homogeneous pathological entity. Furthermore, there is as yet no known normal counterpart with which the hairy cell can be compared.

**Malignant lymphoma of germinal centre cell origin**

Immunological as well as morphological light and
Fig. 10  Hairy cell leukaemia. A portion of hairy cell, which shows finger-like microvilli, three ribosome-lamella complexes, two lysosomal inclusions, and a lipid droplet. (× 46 000)

Fig. 11  Hairy cell leukaemia. A hairy cell with a kidney-shaped pale nucleus, a ribosome-lamella complex (RL), a well-developed Golgi apparatus, and a large pseudopod (P). (× 14 000)
electron microscopic studies (Sordat et al., 1970) indicate that lymphatic follicles and their germinal centres are, at least in their early phase, the site of multiplication of B-lymphocyte clones, which react specifically to antigenic stimulation. In fact, large deposits of immunoglobulin are accumulated within germinal centres (Nossal et al., 1968). Cytologically, in secondary stimulated follicles, several cell types are identifiable. There are specific large cells named germinoblasts (Lennert, 1957; Lennert and Remmele, 1958), renamed centroblasts (Gerard-Marchant et al., 1974), and small cells named germinocytes (Lennert, 1964) renamed centrocytes 1, and, in addition, dendritic reticulum cells (Milanesi, 1965a, 1965b), 'tingible body' macrophages (histiocytic reticulum cells), a few plasma cells and precursors, and occasional immunoblasts (ie, large basophilic transformed lymphocytes). In germinal centres mitotic activity is always very intense.

Normal centroblasts are round cells measuring between 12 and 15 μ with basophilic cytoplasm and a rather clear nucleus in which two to three nucleoli are located against the nuclear membrane. Normal centrocytes are small cells with little cytoplasm, an indented nucleus, and very small nucleoli. These cells are easily identified in thin and well-stained sections of properly fixed material. They are also readily identified in imprints. The relationship between centroblasts and centrocytes is interpreted as the transformation of the former into the latter on the basis of cytophotometric and autoradiographic investigations which showed that the cellular proliferation in germinal centres is primarily a function of centroblasts (Lennert et al., 1969). Opposing views have been expressed by Lukes and Collins (1974). It appears, however, that non-stimulated primary follicles do not contain centroblasts and are made up predominantly of centrocytes, some of which carry natural antibody on the cell surface. It has been stressed that centroblasts and centrocytes are specific transformed lymphoid cells of the germinal centres which have unique distinctive properties. It has also been emphasised that it seems inadvisable to use for their identification inappropriate names, such as histio-
cytes or poorly differentiated lymphocytes (Lennert, 1973). It also seems unrealistic to define the cells of germinal centres by use of the recommended terms, such as large lymphoid cells and medium-sized lymphocytes (Cottier et al., 1973), since the same are also applied to other cell types seen in histological sections of lymph nodes that have different staining properties of the cytoplasm, structure of the nucleus, and size and number of nucleoli.

The derivation of ML from the cells of the germinal centres has been the subject of a long-standing debate. Morphological light microscopic and ultrastructural data (Lennert et al., 1966; Mori and Lennert, 1969; Kojima et al., 1973; Lennert, 1973; Kaiserling, 1977a), immunological findings (Shevach et al., 1973; Jaffe et al., 1974a), and cytochemical data (Lennert, 1964, 1968) were accumulated essentially in favour of a derivation of ML from the cells that are the constituents of the primary and secondary lymphatic follicles (Lennert et al., 1975).

CENTROCYTIC MALIGNANT LYMPHOMA (GERMINOCYTOMA)

Lymphomas made up of cells that correspond to the normal centrocyte have also been defined as poorly differentiated, lymphocytic ML. In lymph nodes, these tumours always reveal a diffuse growth pattern, even though they may also display a vaguely nodular aspect (Fig. 12). In such cases there are ill-defined, small, compact nodules, which closely resemble primary follicles, but a definite follicular architecture with compression of the periphery of the reticulin fibre network and of the vessels is never seen. The centrocytes reveal a rather wide range of sizes. In general, they are small or medium-sized (Kaiserling, 1977a), have cleaved and indented nuclei, usually with one or rarely two very small nucleoli that are distant from the nuclear membrane, and a small amount of slightly basophilic PAS-negative cytoplasm. The main differences from lymphocytes of B-CLL in sections consist of the slightly larger size, the nuclear indentations, and the chromatin structure; however, the nuclear irregularities are less impressive in imprints. In cases in which the cells are predominantly small and not heavily indented, the diagnosis of lymphocytic intermediate type ML has been applied (Dick et al., 1974). However, even small indented cells are different from the small lymphocytes of CLL, as has also been shown in viable cells (Schrek and Donnelly, 1961).

There are cases in which the centrocytes are quite large, with a size that is double that of the small ones. ML made up of large centrocytes is a separate subgroup termed large, cleaved, follicular centre cell ML of Lukes and Collins’ (1974) classification. Large centrocytes have prominent nucleoli and a basophilic cytoplasm, show marked atypia, and reveal a high mitotic index. Cell kinetic studies indicate that large cell centrocytic ML has a higher degree of proliferation than do the small cell types (Silvestrini et al., 1977). For these cases centrocytic sarcoma was the diagnosis proposed by Kaiserling (1977a).

Of the accompanying cells centroblasts are usually recognisable in sections but centroblast-like cells may

1Greek: χιντρον = point, centre; βλαστῶν = offspring, germ; χύτων = cavity, container.
Malignant lymphoma, centrocytic: axillary lymph node, 59-year-old man. Small-sized centrocytes with indented nuclei and small nucleoli. Possible dendritic cells (arrow). Diffuse type of growth. (Giemsa × 400)

Malignant lymphoma, centroblastic-centrocytic, diffuse: mesenteric lymph node, 38-year-old man. The two cell types are approximately equally represented. (PAS × 400)

Malignant lymphoma, centroblastic: mesenteric lymph node, 27-year-old man. Monomorphic diffuse growth of centroblasts. (Giemsa × 400)
occasionally be identified in imprint preparations of affected lymph nodes. Furthermore, a small number of plasma cells and precursors may be found. In some cases there is a variable amount of sclerosis. Interspersed dendritic reticulum cells are a frequent finding even on light microscopy which is confirmed on electron microscopy (Lennert et al., 1975b). In imprints of lymph nodes and smears of extranodal centrocytic ML, centrocytes are quite typical because of their nuclear shape and the lightly basophilic, small cytoplasm (Rilke et al., 1978b). Small cell centrocytic ML seems to be among those tumours the cells of which have been shown to have immunological properties intermediate between CLL and follicular centroblastic-centrocytic ML (Jaffe et al., 1977) and frequently to show a strong alkaline phosphatase activity on their cellular membranes, thus reflecting similarities to the cells of primary follicles (Nanba et al., 1977a). In imprints, centrocytes reveal a moderate acid phosphatase reaction and a cell membrane-bound ATPase activity (Kaiserling, 1977a).

CENTROBLASTIC-CENTROCYTIC MALIGNANT LYMPHOMA (GERMINOBLASTOMA)

Even if Brill et al. (1925) and Symmers (1927) are commonly given credit for the first description of follicular ML, the first clear documentation of a ML with a germinal centre-like structure was given in 1916 by Ghon and Roman. This entity, which is practically unknown in childhood (Butler, 1969; Lennert, 1973; Hausner et al., 1977), is most commonly, at least in the initial stages, made up of follicular structures of variable size, which are very likely to be the malignant counterpart of the secondary lymphatic follicle, since they contain two proliferating cell types which are present in the normal follicles in various proportions, depending on their functional activity (Lennert and Müller-Hermelink, 1975). De novo diffuse centroblastic-centrocytic ML is uncommon, while in a number of cases follicular and diffuse patterns are present at the same time. Cytologically, the neoplastic population is made up of centrocytes, which are identical with those described previously, and of centroblasts which may also show some variation in size. Their nucleus is round and contains a delicate chromatin network with two or three prominent nucleoli, which are adjacent to the nuclear membrane. The cytoplasm is basophilic and pyroninophilic and does not contain PAS-positive material. In imprints, centrocytes and centroblasts reveal a moderate acid phosphatase activity. 5'-Nucleotidase and adenosinenucleophosphatase activities were found at the cellular membrane of centroblasts and centrocytes (Lennert and Rinnerberg, 1961; Müller-Hermelink, 1974; Kaiserling, 1977a). A variant of this lymphoma with a variable degree of sclerosis was described by Bennett and Millett (1969) and has a better prognosis (Bennett, 1975).

Proteinaceous, eosinophilic, and PAS-positive precipitate is present in some of the neoplastic follicles (Rosas-Urble et al., 1973). Intercellular accumulation of immunoglobulin and/or antigen-antibody complexes in neoplastic follicles is much less evident than it is in benign hyperplastic germinal centres (Braylan and Rappaport, 1973). The presence of dendritic reticulum cells may already be suspected on light microscopy in sections and in smears, but it can be better demonstrated ultrastructurally (Lennert and Niedorf, 1969; Glick et al., 1975; Lennert and Müller-Hermelink, 1975; Levine and Dorfman, 1975). When the centrocytes predominate, as is more commonly the case, the histological diagnosis of nodular, poorly differentiated lymphocytic ML is usually applied. When centrocytes and centroblasts are present in a similar proportion, then the diagnosis of nodular mixed lymphocytic and histiocytic ML is usually made, whereas when the large cells predominate, the diagnosis of nodular histiocytic ML is applied. The most common combinations are the first and the second. The mitotic index is usually low, but it is increased when the centroblasts predominate. Follicular lymphomas may maintain their follicular pattern for a long time, even until the death of the patient (Warnke et al., 1977); however, they may also tend to transform into the diffuse form (Fig. 13), while the reverse has never been observed (Rappaport et al., 1956). The morphological steps of the transformation from the follicular to the diffuse type of growth pattern have already been illustrated (Rappaport et al., 1956; Rappaport, 1966; Lukes and Collins, 1975). The evolution to the diffuse growth may either not produce changes in the cellular population or accompany an abrupt increase either of centroblasts or, rarely, of anaplastic centrocytes. The increase of the centroblastic component is paralleled by the increase of the cellular proliferation rate, as has been shown by cell kinetic studies (Silvestrini et al., 1977). Centroblastic-centrocytic ML may eventually undergo an anaplastic transformation, and in that case identification of the histological type becomes difficult. Differential diagnosis between centrocytic and centroblastic-centrocytic, diffuse ML has been discussed in detail (Kaiserling, 1977a).

Malignant lymphomas made up exclusively or almost exclusively of cells identifiable as malignant centroblasts, that is, large, round cells with basophilic cytoplasm and a vesicular nucleus with two or three nucleoli adjacent or adherent to the nucleus membrane (Fig. 14), are usually diagnosed as histo-
Fig. 15  Malignant lymphoma, centrocytic. Two centrocytes showing deeply indented nuclei with condensed chromatin and small nucleoli. (× 10,000)

Fig. 16  Malignant lymphoma, centroblastic-centrocytic, diffuse. A centrocyte with nuclear blebs. (× 11,000)

Fig. 17  Malignant lymphoma, centrocytic, large cell subtype. The large centrocytes have an indented nucleus with diffuse chromatin. (× 10,000) Inset: the Golgi apparatus is more developed than in the usual centrocytes. (× 22,000)
centrocytic ML. In sequential biopsies of centroblastic-centrocytic ML and at necropsy (Lennert, 1976) of former centroblastic-centrocytic ML, the transformation into a pure centroblastic ML (germinoblastic sarcoma) may become evident and should be considered as the transition to a high-grade ML with a poor prognosis. Although infrequently, primary centroblastic malignant lymphomas do exist as de novo malignancies and may occasionally be follicular (Kaiserling, 1977a), even if more often they appear with a diffuse growth pattern. The most prominent feature of centroblastic ML is the uniform proliferation of highly malignant-looking cells, which retain the morphological and immunological characteristics of the centroblast. Cytologically, these cells are larger and different from those of Burkitt type ML, in which the nucleoli are more variable in number and size and commonly not adherent to the nuclear membrane and the cytoplasm is more deeply basophilic and contains lipidic droplets. Quite often the proliferating cell population is also made up, in addition to the malignant centroblasts, of some monomasts and centrocytes. Immunological studies on six cases of large cell ML, which developed in patients with a previous diagnosis of 'nodular' lymphoma, revealed the persistence of B-lymphocyte markers (Jaffe et al., 1977).

Ultrastructurally, the centrocytes have a diameter (5-10 μ) and cytoplasmic characteristics that approach those of the small circulating lymphocytes, from which they differ mainly in the morphology of their nuclei. Centrocytes (Fig. 15) have a deeply cleaved nuclear membrane, which confers on the nucleus a characteristic lobed appearance with lobes connected by narrow bridges. Heterochromatin is less dense than it is in normal small lymphocytes, and nucleoli are inconspicuous or absent. Nucleolar pockets (blebs) are sometimes found (Fig. 16). The cytoplasmic areas surrounded by the nuclear pockets occasionally contain strictly interlaced tubules of smooth endoplasmic reticulum (Fig. 16). Their scanty cytoplasm contains a few small mitochondria and rough endoplasmic reticulum cisternae, a small Golgi apparatus, and numerous monoribosomes. Centrioles, lipid droplets, small microfilament bundles, and lysosome-like dense granules are observed in some cells. Neoplastic centrocytes are similar to normal cells but their nuclei are more wrinkled and show a larger number of blebs than do those of normal cells.

Large centrocytes (Fig. 17) have similar twisted nuclei but are larger (7-14 μ in diameter) and show more signs of morphological activation than do the small centrocytes. The chromatin is more dispersed, the nucleoli, the Golgi apparatus (Fig. 17, inset), and the rough endoplasmic reticulum are more prominent, and polyribosomes and mitochondria are more abundant than they are in small centrocytes.

The centroblasts (Fig. 18) measure 7 to 30 μ in diameter, have large oval or oblong nuclei, dispersed chromatin, and one or more prominent nucleoli with a well-developed nucleolonema. Nucleoli are either central or placed against the nuclear membrane. Although the nuclei of centroblasts may show some irregularities, these are not as conspicuous as they are in centrocytes. Nuclear blebs are seldom observed. Polyribosomes are much more abundant and larger than they are in centrocytes, while monoribosomes are rarely present in the cytoplasm. Several electron-lucent mitochondria, a well-developed Golgi apparatus, a small amount of granular and agranular endoplasmic reticulum, and a few lysosomes are also found in the cytoplasm. Nuclei of neoplastic centroblasts occasionally display more indentations and irregularities than do those of normal cells.

In a few cases of centroblastic-centrocytic ML, both tumour cell types (Figs 19, 20), and especially the centrocytes, have a well-developed rough endoplasmic reticulum (Kaiserling, 1977a). The long and flat cisternae either assume a circular arrangement around the nucleus (Fig. 20) or form whirl-like structures. The cells with a well-developed endoplasmic reticulum also present more numerous mitochondria and lysosome-like dense bodies (Fig. 20, inset) and a better developed Golgi apparatus than do the usual centroblasts and centrocytes.

Dendritic cells are easily observed among tumour cells in follicular centroblastic-centrocytic ML. Such cells are less common in the diffuse variant (Levine and Dorfman, 1975). The dendritic cell plasma membrane forms long, branch-like processes, which sometimes interdigitate with those of other dendritic cells. Junctional apparatuses of the macula adherens type are observed at the sites of membrane contact (Fig. 21). Numerous fine cytoplasmic filaments condense, especially in connection with desmosomes. Phagocytosis is absent. The dendritic cell nucleus has a characteristic wavy shape. Chromatin is diffusely dispersed and only slightly condensed along the nuclear membrane. Nucleoli have a well-developed nucleolonema. An inconspicuous Golgi apparatus, scattered mono- and polyribosomes, a few mitochondria and rough endoplasmic reticulum cisternae, and rare ribosomes are present in the cytoplasm.

The concept that the ML described above are actually neoplasms of germinal centre cells has been strongly supported by immunological studies. Complement receptors were reported to be characteristic of B cells within lymphoid follicles (Braylan et al., 1975; Jaffe et al., 1974a, 1975), while surface
Morphology of lymphatic cells and of their derived tumours

Fig. 18 Malignant lymphoma, centroblastic-centrocytic. A centroblast presenting a roughly oval nucleus with dispersed chromatin and three well-developed nucleoli adjacent to the nuclear membrane. (× 9000)

Fig. 19 Malignant lymphoma, centroblastic-centrocytic, diffuse. A centroblast with well-developed rough endoplasmic reticulum. (× 12000)

Fig. 20 Malignant lymphoma, centroblastic-centrocytic, diffuse. A centrocyte with abundant flat endoplasmic reticulum cisternae and some lipid droplets. (× 12000) Inset: numerous lysosome-like dense bodies in a centrocyte with abundant rough endoplasmic reticulum. (× 9000)
immunoglobulin-bearing B cells are distributed in both follicles and the medullary cords. It was shown that follicular lymphomas are of B-cell type origin and that all their cells have complement receptors and the major bear surface immunoglobulin (Gajl-Peczalska et al., 1973; Shevach et al., 1973; Jaffe et al., 1974a; Aisenberg and Long, 1975; Leech et al., 1975a; Bloomfield et al., 1976). The same applies to centrocytic ML (Stein, 1975). The cells of follicular lymphomas have neither receptors for cytophilic antibody nor do they form spontaneous E rosettes. The presence of the complement receptor is unrelated to the cytological subtype of follicular lymphoma, although a loss of these receptors was reported during the transition from the nodular to the diffuse pattern (Crossman et al., 1977). Except for a few IgG-containing cells, none of the tumour cells contains cytoplasmic immunoglobulin (Johansson et al., 1976). In contrast to B-CLL lymphocytes, the small follicular centre cells of cases with leukaemic manifestations and either a nodular or diffuse structure show a lesser tendency to form rosettes with mouse erythrocytes (Koziner et al., 1977). Quite conceivably, the cells of diffuse centroblastic-centrocytic ML with anaplasia may behave as 'receptor-silent' cells or show immature receptor profiles, as has been reported (Habeshaw et al., 1977) for some diffuse, mixed, and histiocytic ML, which correspond to large cleaved and non-cleaved follicular centre cell ML described by Lukes and Collins (1974).

It has been postulated that for the characterisation of centroblastic-centrocytic ML, further investigation of the tissue between the neoplastic follicles may yield significant additional information. In fact, morphologically on light and electron microscopy, the interfollicular tissue contains many structures (postcapillary venules with recirculating lymphocytes, T-associated plasma cells, interdigitating reticulum cells) that are characteristic of the paracortical area of lymph nodes (Kaiserling, 1977a). In addition, immunologically, at the periphery of the neoplastic follicles, an unforeseen accumulation of T lymphocytes has been detected (Jaffe et al., 1977). Whether this finding should be interpreted as a host defence or as a process relevant to the follicular tumorigenesis remains a challenging question.

In a number of cases, ML of germinal centre cell origin presents with leukaemic spread, the first description of which may be identified in the report of Isaacs (1937). He reported a type of lymphocytic leukaemia associated with lymphosarcoma in the lymph nodes, which he believed to be different from other types of lymphocytic leukaemia. Lymphosarcoma cell leukaemia may occur either as an early phenomenon (early leukaemic lymphosarcoma) or may appear as a late manifestation (late leukaemic lymphosarcoma) (Mathé et al., 1976). In all cases of blood involvement the marrow is invariably infiltrated, but whether or not marrow infiltration without blood involvement should be considered leukaemic conversion is debatable (Wintrobe et al., 1974).

Owing to the use of the Rappaport classification, in which nodular, poorly differentiated lymphocytic ML comprise pseudonodular centrocytic as well as follicular centroblastic-centrocytic ML, and diffuse, poorly differentiated lymphocytic ML comprise centrocytic and centroblastic-centrocytic diffuse ML, it is difficult to reinterpret previous reports, since quite often nodular and diffuse ML cases are reported together. The impression that one gains, however, is that in both instances centrocytes are the most frequently circulating cell type (Figs 22 and 23). In any case, centrocytes in blood smears and in marrow smears bear striking similarities to the 'notched-nucleus' cells described by Anday and Schmitz (1952) in follicular lymphomas, to the 'haemato-gones' described by Rosenthal et al. (1952) also in follicular lymphomas, and to the cells observed in some of the cases described by Schwartz et al. (1965). The same cell type corresponds to the third type of leukaemic lymphosarcoma described by Mathé et al. (1975b) and to the notched-nucleus lymphocytes described by Spiro et al. (1975). The cytoplasm may contain vacuoles, azurophilic granules, and basophilic granules, which appear structureless on electron microscopy (Wintrobe et al., 1974). In a series of 16 cases with lymphosarcoma cell leukaemia (Schnitzer et al., 1970) lymph node biopsy revealed a poorly differentiated lymphocytic ML, which was diffuse in six and nodular in 10 cases. In the majority of diffuse ML leukaemia was present at diagnosis but was a later manifestation in the nodular cases. The changes in the course of the disease were also demonstrated by the fact that, at the time of necropsy, one case only had maintained the nodular structure of the lymphoma, while the others had changed towards a diffuse pattern.

In the blood of centroblastic-centrocytic follicular ML, mostly centrocytes are recognisable, whereas centroblasts may be found only occasionally (Lennert, 1969). The circulating centrocytes reveal the same previously described immunological surface markers as do the cells in the solid tumour, thus confirming their B-type nature and germinal centre origin. There are, however, reports on dual markers for B and T cells detected in cases of lymphosarcoma cell leukaemia (Hsu et al., 1975; Lin and Hsu, 1976).

In a survey of 75 cases of follicular lymphoma, among which all cytological subtypes (small and large cells) were represented, 25 cases were found to
have abnormal cells in the blood (Spiro et al., 1975). In 17 the blood lymphocyte count ranged between 5 and 30 × 10^9/l, and in another eight cases it was below 5 × 10^9/l; in all these cases, however, the marrow aspirate films showed neoplastic lymphocytic infiltration made up of abnormal notched-nucleus cells. These were also found in blood films of cases of nodular lymphoma with large cell type predomina-
ice (histiocytic). The most important defining characteristic of the circulating cells, in addition to the nuclear clefts of varying depth, was the chromatin pattern, which was more homogeneous than was that of the small lymphocytes and did not reveal any structure made up of coarse blocks of densely stained chromocentres. In addition to the cells with the indented nuclei, other circulating cells with scanty basophilic cytoplasm and barely perceptible nuclear notches were identified in the blood.

Examination of marrow trephine biopsy sections at the time of initial diagnosis of follicular lymphomas reveals a high percentage with involvement (Dick et al., 1974; McKenna et al., 1975; Castellani et al., 1977), although the degree of infiltration and the pattern of distribution is variable (Jones et al., 1972, 1973; Dick et al., 1974). Marrow films are somewhat less reliable. For diffuse, poorly differenti-
tated lymphocytic ML, marrow invasion was found by trephine in about 30% of cases. Marrow involvement by nodular lymphomas with predo-
nantly small cells (nodular lymphocytic, poorly differenti-
tated) was much more frequently encountered than in those with large cells (nodular histiocytic) (Jones et al., 1972, 1973).

With regard to the cell types encountered in marrow invasion, the morphological description of a small cell (7-10 μ in diameter) with an irregularly cleaved nucleus, small inconspicuous nucleoli, and scanty, lightly basophilic cytoplasm corresponds to that of a centrocyte, small cleaved follicular centre cell (Lukes and Collins, 1974), small nodular lymphoma cell (McKenna et al., 1975), 'haemato-
gone' (Rosenthal et al., 1952), and the lymphocytic cell of poorly differentiated lymphocytic ML (Dick et al., 1974). If this is the cell most commonly found in these instances, it is also not uncommon to identify both large cells (which measure up to 20 μ in diameter with a round nucleus, multiple nucleoli, and basophilic cytoplasm) that correspond to centro-
blasts and cells with features intermediate between the centrocytes and the centroblasts (Dick et al., 1974; McKenna et al., 1975). The rare circulating large cells are cytologically different from immuno-
blasts inasmuch as their cytoplasm is less baso-
philic and their large nuclei contain multiple nucleoli rather than a single huge central nucleolus.

The morphological differences between haemato-
gones found in the marrow of children and adults in health and in various pathological conditions and the neoplastic haematogones (Rosenthal et al., 1952) have been noted by McKenna et al. (1975). On the other hand, the occasional finding of cells similar to small centrocytes in marrow films in various non-neoplastic haematological conditions has also been reported (McKenna et al., 1975). It seems, therefore, that this cell type is sufficiently distinct to be differentiated on cytological grounds alone from the small lympho-
cyte.

In summary, almost all the reports on the leu-
kaemic manifestations of poorly differentiated lymphocytic ML, both nodular and diffuse, and of nodular ML of other cell types clearly indicate that the haematological, morphological, and immunological features observed are different from those of CLL. It appears that in these cases the circulating cells are exclusively or predominantly centrocytes, which may reveal more or less accentuated features of atypia, whereas the presence of centroblasts is much less common. The need for a better correlation in the future between histological and haematological cell morphology of leukaemic, poorly differentiated lymphocytic ML is justified, since immunological differences between diffuse and nodular types of ML have been reported (Aisenberg and Long, 1975; Aisenberg and Wilkes, 1976). In fact, clinical and haematological differences between the purely centro-
cytic diffuse and the follicular centroblastic-centro-
cytic ML do exist. The greater tendency towards generalised dissemination and marrow invasion (Castellani et al., 1977), and leukaemic change in some cases with a high WBC and a fulminant course, in addition to the existence of the large cell variant (Silvestrini et al., 1977), are some of the distinguishing features of centrocytic ML versus follicular centroblastic-centrocytic ML, which in turn are more frequently associated with a chronic leukaemic manifestation. At the same time it is fair to add that the importance of these distinctions has been doubted (Dorfman, 1977).

BURKITT LYMPHOMA

In spite of clinical (Levine et al., 1975), epidemi-
ological, immunological (Epstein et al., 1976), and virological (Zur Hausen, 1975; Andersson et al., 1976; Ziegler et al., 1976) differences between endemic African (Burkitt, 1958) and sporadic African and non-African (O'Conor et al., 1965; Burkitt, 1967) Burkitt lymphomas, their morpho-
logical-cytological expressions are very similar (Berard et al., 1969). This tumour has been con-
considered as the most classic example of an undifferentiated ML because of its uniform and
monomorphic cellular population without evidence of maturation. In section, the cells measure 7 to 12 μ and have roundish or oval nuclei, which contain two to five prominent basophilic nucleoli, usually not adherent to the nuclear membrane. Abnormalities of the nuclear membrane, such as indentations, are commonly absent. The chromatin is usually made up of large, coarse chromocentres, and the mitotic index is high. The cytoplasm is amphophilic and pyroninophilic and contains small vacuoles.

In imprints of tumour tissue, the cells measure 10 to 25 μ in diameter and have a deeply basophilic cytoplasm. The cytoplasm is usually not abundant and contains droplets of neutral Oil Red O-positive lipids, which correspond to well-defined clear vacuoles in alcohol-fixed preparations.

Furthermore, the cytoplasm usually does not contain PAS-positive material, except for rare coarse granules, and does not reveal any non-specific esterase of peroxidase activity. Small amounts of acid phosphatase may be present. Alkaline phosphatase activity could be demonstrated in some cells (Nanba et al., 1977a). Macrophages reveal a high content of acid phosphatase and non-specific esterase.

A common but not constant phenomenon is the large number of 'tingible body' macrophages similar to those of the secondary follicle among the tumour cells. These are the cells whose relative pallor contrasts in sections with the dark, nuclear-dense masses of lymphoma cells to produce the well-known but non-specific 'starry-sky' appearance. The macrophages contain cellular debris and ingested tumour cells. Dendritic reticulum cells, however, are absent (Kaiserling, 1977a).

Electron microscopically, the picture is also dominated by large lymphoid blast cells mixed with a varying number of non-malignant histiocytes (Epstein and Barr, 1965). In the blast cells the nuclear:cytoplasmic ratio is high (Epstein and Achong, 1965; Flandrin et al., 1975). The pale nuclei are round or oval and occasionally slightly indented. Heterochromatin is scarce and placed along the nuclear membrane. Nuclear blebs are frequently observed. Nucleoli are prominent. The cytoplasm contains a few endoplasmic reticulum cisternae and a characteristically large number of free ribosomes and polyribosomes. Well-developed polarised mitochondria, lipid droplets, probably made up of hydrophobic unsaturated lipids (Feremans et al., 1976), and clear vesicles are also found. The Golgi apparatus is usually inconspicuous but occasionally is relatively well developed. Annulate lamellae are observed in some cases (Epstein and Achong, 1965). The plasma membrane is generally smooth.

The cells of Burkitt lymphoma have been considered to derive from germinal centre cells (small centroblasts) because of their similarities to the normal counterpart within the lymphatic germinal follicle (Lukes and Collins, 1974). Immunologically, in many cases of both African endemic and non-African sporadic Burkitt lymphoma, monoclonal surface-bound immunoglobulin was identified on the malignant cells (Klein, 1971). Tumour cells have been shown to synthesise immunoglobulin also in tissue cultures (van Furth et al., 1972). The same finding, that is, the presence of surface IgM (and occasionally of IgG) with λ light chains, was demonstrated in neoplastic leukaemic cells of non-endemic American Burkitt lymphomas (Flandrin et al., 1975; Mann et al., 1976). In addition, complement receptors (Nussenzweig et al., 1971; Shevach et al., 1972; Stein, 1975) and receptors for aggregated IgG (Flandrin et al., 1975) were identified on a number of cells. Burkitt cells do not form spontaneous E rosettes (Flandrin et al., 1975). Differences in surface marker patterns were reported for endemic Epstein-Barr virus (EBV) genome-containing and for sporadic EBV-negative Burkitt ML (Epstein et al., 1976).

The most relevant recent morphological finding in biopsies and necropsies is the involvement of germinal centres of lymph nodes and Peyer's patches (Mann et al., 1976). In particular, the confinement of foci of Burkitt ML to single germinal centres further supports the suggestion that this lymphoma should be considered a tumour of germinal centre origin. Some remaining uncertainties on the nature of the proliferating tumour cells, their 'lymphoblastic' (that is, immature lymphoid) appearance, the prevalence of the tumour in childhood, and its occasional acute leukaemic manifestation suggested its inclusion among high-grade lymphoblastic ML of the Kiel classification.

Marrow invasion (Wright, 1968) is quite variable; it develops in about 10 to 20% of cases in Africa and is even more common outside Africa (Levine et al., 1975; Brearley et al., 1977). No correlation between the percentage of lymphoma cells in marrow or blood was found in non-African cases (Brunning et al., 1977). The leukaemic manifestation may be present at onset (Bluming et al., 1972) or may develop terminally. The terminal leukaemic pattern is quite rare in Africa (Clift et al., 1963), and its morphology is that of an ALL of B-cell type (Chessells et al., 1977). Outside Africa, Burkitt type ALL has been reported to represent 2% of all ALL (Flandrin et al., 1975). In African (Manolov and Manolova, 1972) and non-African (Hübner and Littlefield, 1975; Philip et al., 1977) Burkitt ML, structural chromosomal abnormalities were found.
Fig. 21 Malignant lymphoma, centroblastic-centrocytic, follicular, and diffuse. A junctional apparatus between the process of two dendritic reticulum cells. (× 51 000)

Fig. 22 Malignant lymphoma, centrocytic. Leukaemic blood smear: WBC 26·5 × 10⁹/l. Circulating centrocytes 26%. Two atypical centrocytes. (May Grünwald-Giemsa × 1000)

Fig. 23 Malignant lymphoma, centroblastic-centrocytic. Leukaemic blood smear: WBC 21·7 × 10⁹/l. Circulating centrocytes 95%. An atypical centrocyte. (May Grünwald-Giemsa × 1000)

Fig. 24 Malignant lymphoma, lymphoplasmacytoid: axillary lymph node, 60-year-old woman. Diffuse growth of small round cells with scattered PAS-positive Russell bodies (arrow). (PAS × 400)

Fig. 25 Malignant lymphoma, lymphoplasmacytoid polymorhous: cervical lymph node, 47-year-old man. Numerous variably sized Russell bodies. (PAS × 400) (Courtesy of Professor L. Tropeano, Rho).
Malignant lymphoma of immunoglobulin-secreting cells and lymphoplasmacytic dyscrasias

LYMPHOPLASMACYTOID MALIGNANT LYMPHOMA

The existence of ML in which a variable proportion of the neoplastic cells show mature or immature plasmacytic or plasmacytoid features is fully recognised. While in some classifications of ML it has simply been stated that some subtypes may show plasmacytoid ‘differentiation’ without therefore creating new subtypes (Bennett et al., 1974), in other schemes well-defined entities are recognised in which the mature and/or immature plasma cell component is the qualifying feature of the neoplasm (Dorfman, 1974; Gerard-Marchant et al., 1974; Lukes and Collins, 1974). In this context it should be noted that discrepancies between morphology and function are not uncommon; one example is primary macroglobulinaemia (Waldenström, 1944), a fairly well-defined clinical entity, which does not correspond to a similarly well-defined morphological counterpart (Stein et al., 1972).

Primary macroglobulinaemia is a progressive and systemic lymphoproliferative disease, which is associated with monoclonal IgM production. In the proliferating cells μκ and/or μλ immunoglobulin was identified, even in paraffin sections, by the immunoperoxidase technique (Pinkus and Said, 1977). The morphological heterogeneity of primary macroglobulinaemia is visualised by the fact that in many cases the predominant proliferating cell type is either the small lymphocyte (Pangalis et al., 1977) or the plasmacytoid lymphocyte (Lennert et al., 1975b; Lukes and Collins, 1975), while in other cases more pleomorphic histological pictures were reported. In addition, some of the patients with this disease later develop large cell lymphoid tumours. On the other hand, Lennert et al. (1975c) and Stein et al. (1974b) reported cases of ML that corresponded to the morphological patterns described above in which macroglobulinaemia could be detected only in a minority of the cases but in which accumulated monoclonal and occasionally polyclonal immunoglobulin could be detected within the tumour tissue. Finally, the identification of ML, still with the same morphology but with production of IgG or IgA, made the existence of Waldenström’s macroglobulinaemia as a pathological entity questionable. From a diagnostic point of view it seems more important to realise that the existence of ML made up of lymphocytes with plasmacytoid features or of mixtures of lymphocytes and plasma cells with increased tissue immunoglobulin should be recognised, regardless of whether or not there is a monoclonal increase of IgM or of other immunoglobulin in the serum.

A number of cases of lymphoplasmacytoid ML have probably been diagnosed in the past as diffuse well-differentiated lymphocytic ML, as was recently pointed out by Pangalis et al. (1977), who found that 90% of patients with this lymphoma associated with monoclonal gammopathy revealed plasma cells and/or plasmacytoid lymphocytes in their neoplasms. The same type of tumour is considered a distinct entity among B-cell type ML that derive from the plasmacytoid lymphocyte (Lukes and Collins, 1974).

The morphological normal counterpart of one of the proliferating cells has been identified by Lennert (Lennert and Müller-Hermelink, 1975) as the lymphatic plasma cell originally described by Moeschlin. This cell is smaller than the classic reticulum or Marschallo (1895) plasma cell, has a smaller Golgi body, and a scanty cytoplasm, and the nucleus is not eccentric. However, the structure of the cytoplasm does not differ qualitatively on electron microscopy from that of the reticulum plasma cell, but on the surface of circulating lymphatic plasma cells IgM could be detected (Lennert and Müller-Hermelink, 1975). It has been postulated that lymphatic plasma cells may originate rapidly from B lymphocytes, bypassing the stage in the follicular centres, and become responsible for IgM synthesis. Reticulum plasma cells, after maturation within the germinal centres (Nieuwenhuis and Keuning, 1974) are responsible for the production of IgG and IgA. Therefore, lymphoplasmacytoid ML were defined also as immunocytomas (Stein et al., 1974b; Lennert et al., 1975c) with reference to the functional nomenclature proposed by Dameshek (1967).

Three main histological variants of lymphoplasmacytoid ML can be recognised (Lennert et al., 1975c; Schwarze et al., 1976) in sections of lymph nodes or in other common extranodal sites, such as the subcutaneous tissue, the orbit, the leptomeninges, and the gastrointestinal tract (Rilke et al., 1978b). The first variant is the lymphoplasmacytic subtype, which is made up of a mixture of small lymphocytes, identical with those of CLL, and of immature as well as mature Marschallo-type plasma cells. The second variant is the lymphoplasmacytoid ML in which, in addition to some lymphocytes, there is a uniform proliferation of lymphatic plasma cells, which are slightly larger than the lymphocytes and have a scanty, homogeneous basophilic cytoplasm. The nucleus is central or slightly eccentric and contains moderately coarse or delicate chromatin and a barely visible central nucleolus. Mast cells are more numerous than they are in CLL, and their number helps in the differential diagnosis in equivocal cases (Satodate et al., 1977). The reticulin fibre network is fairly abundant.
The third variant is the pleomorphic lymphoplasmacytoid ML in which a mixture of lymphocytes, lymphatic plasma cells, centroblasts, centrocytes, and immunoblasts dominates the picture. It has been shown that the degree of pleomorphism of this ML parallels the proliferation rate (Silvestrini et al., 1977). Pleomorphic malignant lymphomatous proliferations with plasmacytic differentiation, polyclonal hyperimmunoglobulinaemia, and autoimmune haemolytic anaemias were reported (Flandrin et al., 1972).

The cytoplasm of the proliferating cells often shows a diastase-resistant, granular PAS positivity, which is related to the glycoprotein content. In addition, in a very variable number of cells there are round globular inclusions both in the cytoplasm (Figs 24 and 25) and in the nuclei that are PAS positive (Kaiserling et al., 1973); they represent accumulated immunoglobulin, mostly IgM but occasionally also IgG or IgA (Stein et al., 1972). In about 30% of these cases, immunoglobulin of the same class in the serum may be increased. The cytoplasmic inclusions have been referred to as Russell's bodies (Russell, 1890) and the intranuclear inclusions as Dutcher's bodies (Dutcher and Fahey, 1959). Intranuclear material seems to derive from extranuclear accumulation of immunoglobulin in the perinuclear cisternae, with subsequent invagination into the nuclear structure (Brunning and Parkin, 1976).

All lymphoplasmacytoid ML show a diffuse proliferation pattern with occasional pseudonodular configuration. Remnants of secondary follicles may persist. Evolution to basophilic large cell immunoblastic ML can occur (Lennert et al., 1975b). Leukaemic manifestation develops in about one-third of the cases and is associated with a moderately high WBC count, which seldom exceeds $30 \times 10^9/l$. The clinical picture of CLL may predominate, particularly in cases that may be considered borderline, such as those reported in a retrospective study on a series of cases of CLL among which a certain percentage was actually characterised by a leukaemic proliferation of lymphoplasmacytoid cells (Rudders, 1976). In marrow and blood smears the same cells can be identified and differentiated from CLL lymphocytes by virtue of their cytoplasmic basophilia and the slightly larger area of the cytoplasm. The leukaemic manifestation of immunocytomas occasionally shows the cytological pattern of plasma cell leukaemia.

The description of a method for the demonstration of specific immunoglobulin in plasma cells and lymphoplasmacytoid cells in formalin-fixed and paraffin-embedded material using a peroxidase-conjugated antibody and an indirect sandwich technique (Taylor and Burns, 1974) has greatly facilitated the identification of the function of these cells in benign and neoplastic conditions. The electron microscopic investigation of a series of lymphoplasmacytoid ML shows a wide spectrum of morphological variations, ranging from a pattern that consists of a large number of lymphocytes and a much smaller number of plasma cells, lymphoplasmacytoid cells, and immunoblasts, to an almost typical plasmacytoma. In some cases centroblasts and centrocytes are also found. Quite often the neoplastic cells are indistinguishable from normal cells.

Lymphoplasmacytoid cells (Fig. 26) have a nucleus with abundant heterochromatin and an inconspicuous nucleolus, similar to the nucleus of the small circulating lymphocytes. In the cytoplasm, more or less abundant long and flat rough endoplasmic reticulum cisternae, placed around the nucleus, are observed (Lennert and Müller-Hermelink, 1975). Plasma cells with the mature appearance of the Marschalko type have a very abundant rough endoplasmic reticulum, a large Golgi apparatus, and nuclear chromatin, which forms clumps lying against the nuclear membrane; they may also have a nucleus with diffuse chromatin and less abundant rough endoplasmic reticulum, indicating a lesser degree of maturation (Fig. 27). It is frequently possible to follow a spectrum of morphological variants from the plasmacytoid cells to the plasma cells. Deeply indented lobed nuclei are frequently observed in plasma cells of some cases of immunocytoma (Fig. 28) and plasma cell leukaemia (Klein et al., 1977).

Immunoblasts have a cytoplasm that contains a scanty endoplasmic reticulum, very numerous polyribosomes, and a well-developed Golgi apparatus. The nucleus is round with dispersed chromatin and large nucleoli (Florentin, 1975).

Intranuclear inclusions, which are bound by a single membrane and which usually contain a granular or fibrillar material (Kuhn, 1967; Mori and Lennert, 1969; Cawley and Hayhoe, 1973), are observed in both plasma cells and lymphoplasmacytoid cells. These inclusions correspond to the PAS-positive Dutcher bodies (Dutcher and Fahey, 1959; Napoli et al., 1977). Russell bodies (Fig. 28) and crystalline inclusions of the rough endoplasmic reticulum are also occasionally observed (Bessis, 1961; Argani and Kipkie, 1965).

As for the possible derivation of lymphoplasmacytoid ML, their relationship to both germinal centre cells and cells that develop outside germinal centres has been considered (Kaiserling, 1977a): in one-half of the cases complement receptors as well as surface immunoglobulin may be detected on the cells, whereas in the other half no complement receptors are
Fig. 26  Malignant lymphoma, lymphoplasmacytoid polymorphous. A lymphoplasmacytoid cell with abundant heterochromatin, an inconspicuous nucleolus, and flat endoplasmic reticulum cisternae placed around the nucleus. (× 12 000)

Fig. 27  Malignant lymphoma, lymphoplasmacytic. An immature plasma cell that shows a nucleus with diffuse chromatin and a small nucleolus. Dense inclusions are observed in the mitochondria. (× 12 000)

Fig. 28  Malignant lymphoma, lymphoplasmacytoid polymorphous. A plasma cell with a lobed nucleus and Russell bodies within the rough endoplasmic reticulum cisternae. (× 11 000)

Fig. 29  Malignant lymphoma, immunoblastic: axillary lymph node, 66-year-old man. Diffuse proliferation of large cells with basophilic cytoplasm and large central nucleolus. A few cells reveal plasmacytoid features. (Giemsa × 400)
found (Stein, 1975). Lymphoplasmacytoid ML have therefore been considered neoplasms of B cells, the range of which goes from cells that are halfway between the non-secreting and the secreting stage to those that are capable of secretory activity but in a majority of the cases are unable to discharge the product into the blood (Stein, 1975). In fact, less than one-third of the cases of lymphoplasmacytoid ML are diagnosed clinically as Waldenström’s disease. Owing to the variable morphological pattern, Waldenström’s disease has also been defined as a lymphoplasmacytic dyscrasia with abnormal secretion of monoclonal immunoglobulin. The same definition may also be applied to heavy chain diseases, which are characterised by the secretion of subunits of immunoglobulin.

**HEAVY CHAIN DISEASES**

Heavy chain diseases comprise a group of conditions in which populations of lymphoid cells synthesise and secrete immunoglobulin heavy chains that are structurally defective (Buxbaum, 1976). They have also been defined as immunoproliferative disorders which are characterised by the presence in the serum of a monoclonal population of molecules composed of incomplete heavy polypeptide chains which belong to a given class or subclass of immunoglobulin and are devoid of light chains (Warner et al., 1974). Three types have been recognised so far: heavy chain disease (Seligmann et al., 1968), gamma chain disease (Franklin et al., 1964), and mu chain disease (Ballard et al., 1970; Forte et al., 1970).

The diagnosis is based exclusively on non-morphological investigations; the clinical pattern is different from that of multiple myeloma and appears to be extremely variable. Even though in some cases no morphological evidence of abnormal cellular proliferation is found, in most cases mixed patterns similar to those found in macroglobulinaemia are seen in marrow smears and in lymph nodes.

Except for a few cases with a respiratory form of the disease (Florin-Christensen et al., 1974), alpha chain disease affects predominantly the digestive tract, producing malabsorption (Rambaud et al., 1968). Even though alpha chain disease may occasionally be associated with immunoblastic ML of the intestine, in the majority of cases it is associated with what has been defined as Mediterranean lymphoma or immunoproliferative small intestinal disease of Mediterranean type (Rambaud and Matuchansky, 1973). There is a diffuse and massive infiltration of the lamina propria of the intestine by plasma cells and/or lymphocytes (Rappaport et al., 1972; Seligmann, 1975) without evidence of cellular atypia. The proliferation may also be present in mesenteric nodes, but it never spreads to either the marrow or to extra-abdominal sites; it usually begins in the upper small intestine and subsequently involves it completely. The cellular composition varies from completely mature plasma cells to a mixture of plasma cells and lymphocytes or to intermediate lymphoplasmacytoid cells. The mature cell proliferation may occasionally be moderately invasive with atrophy of the intestinal villi. In other cases the proliferating plasma cells are more obviously malignant, and they invade the muscular layer of the intestine. Finally, there are cases in which there is a highly malignant proliferation of immunoblasts, which may or may not be associated with a still evident benign lymphoplasmacytoid proliferation (Ala et al., 1976). A word of caution is needed concerning the neoplastic nature of the benign types, since the condition is rarely reversible either spontaneously or after antibiotic treatment with disappearance of the plasma cell excess and of the abnormal alpha chain (Ala et al., 1976). It has been proved that the lymphomas arising late in the course of the intestinal form of alpha chain disease derive from the same B-cell clone as does the initial plasma cell proliferation (Brouet et al., 1975b, 1977; Pangalis and Rappaport, 1977; Ramot et al., 1977). By contrast, ‘Western type’ ML of the small bowel have clinical and morphological differences and seem to be made up of a different cell type (Lewin et al., 1976).

Gamma chain disease often shows intermediate features between multiple myeloma and ML. It has in common with the former anaemia, frequency of infections, and a variable clinical course, often with severe renal disease. However, there are no lytic bone lesions. In common with ML there is often enlargement of lymph nodes, splenomegaly, and hepatomegaly. It may be associated with a lymphoplasmyctotic ML, either leukaemic or non-leukaemic, or with pleomorphic cells in transition to immunoblastic ML. It may occasionally be associated with extra-osseous plasmacytoma and may show plasma cell leukaemia in the terminal phase.

Mu chain disease is rare and has been sometimes found in association with an immunocytoma (Lennert, personal communication, 1978). In marrow samples, in addition to lymphocytosis, a certain degree of plasmacytosis is found. The plasma cells have been reported to be vacuolated. In both gamma and mu chain disease, cases without evidence of abnormal cell proliferation have also been reported (Seligmann, 1972; Danon et al., 1975).

**PLASMACYTIC MALIGNANT LYMPHOMA**

Pure plasmacytic ML of extra-osseous sites was added later to the low-grade ML of the Kiel classification (Gerard-Marchant et al., 1974). It is equiva-
lent to the extramedullary plasmacytoma (Wiltshaw, 1976), which may arise in lymph nodes, in the upper and lower air passages and the lung, in the spleen, in the skin and subcutaneous tissues, in the gastro-intestinal tract (stomach, small intestine, large intestine), and less frequently in other sites. It has been shown that extramedullary plasmacytoma is composed of an autonomous and irreversible proliferation of plasma cells of varying degrees of differentiation and has characteristic features, which distinguish it from myelomatosis and from solitary myeloma of the bone (Wiltshaw, 1976). The tumour masses are made up of mature plasma cells of the Marschalko reticular type with an eccentric nucleus and typical arrangement of the chromatin; the nucleus is surrounded by a clear halo, and the cytoplasm is intensely eosinophilic and pyroninophilic and shows diastase-resistant PAS positivity. In addition, less well-differentiated and poorly differentiated plasma cells may be present to a variable extent, and in some cases the tumour is made up of very pleomorphic plasma cells. In these the nuclei are more centrally located and may be multiple, the nucleolus is more prominent, and the cytoplasm is more scarce; however, the perinuclear halo is often still present.

In lymph nodes the plasmacytic ML may be either the primary or the metastasis of an extramedullary plasmacytoma (Fishkin and Spiegelberg, 1976). Extranodal extramedullary plasmacytomas show a high incidence of metastatic spread to soft tissues (Wiltshaw, 1976). It has been pointed out that in spite of the anatomical distribution of plasma cells, which is similar to that of lymphocytes, plasmacytic tumours are rare in lymph nodes and much more common in the bone marrow. An unsuspected proportion of plasma cell tumours among the ML arising in the gastrointestinal tract has been reported recently (Henry and Farrer-Brown, 1977). At their presentation, extramedullary plasmacytic ML are rarely associated with serum protein abnormalities, whereas at a later stage paraproteinaemia may appear (Wiltshaw, 1976).

**Immunoblastic malignant lymphoma**

The identification of this entity among the heterogeneous group of more or less differentiated reticulum cell sarcomas (Oberling, 1928; Roulet, 1930; Robb-Smith, 1938; Rössle, 1939) depends upon electron microscopy and immunological methods (Kaiserling et al., 1973; Stein et al., 1972, 1974a). Immunoblastic ML is characterised by a monomorphic proliferation of large cells (15-30 μ in diameter) with pale eosinophilic but deeply pyroninophilic cytoplasm, a moderate or large amount of endoplasmic reticulum, round or oval clear nuclei, one or two large, centrally located eosinophilic nucleoli, and, in some cases, the presence of condensed IgM in the cytoplasm. Mitotic figures are common. The similarities of the neoplastic cells to transformed lymphocytes led to the introduction of the term immunoblastic ML derived from the original terminology of Dameshek (1963).

These neoplasms have previously been classified either as undifferentiated non-Burkitt ML or as diffuse poorly differentiated histiocytic ML. Immunoblastic ML are high-grade malignancies and have a very poor prognosis, allowing only a short survival of the patient. Although in many instances the neoplasm appears rapidly de novo, in a number of cases it represents the late transformation of lymphoplasmacytoid ML, of immunoblastic lymphadenopathy (Lukes and Tindle, 1975), of plasma cell proliferations of various types, and even of follicular lymphomas (Dick et al., 1977). In these latter cases the B-cell type origin of the tumour is very likely, since the common clonal origin of the lymphoplasmacytic proliferation in alpha chain disease and of immunoblasts in the supervening ML could be ascertained.

The cytoplasm contains diastase-resistant PAS-positive granules but no naphthol esterase activity (Stein et al., 1972; Glick et al., 1975), or acid phosphatase (except for a few granules), whereas the enzymatic activities are highly evident in phagocytes containing reticulum cells (Carr, 1973). Membrane-bound ATPase activity is strong, while 5'-nucleotidase is absent (Kaiserling, 1977a).

The identification of these cells in imprints of lymph nodes, as well as in smears of blood and marrow, is based upon the above-mentioned features, and differential diagnosis has to take into consideration Burkitt type ML, centroblastic ML, true histiocytic neoplastic proliferations, and myeloid sarcoma. In particular, in haematoxylin-eosin-stained sections, the cells of the latter may be difficult to identify as myeloid in nature unless there are myelocytes and eosinophilic cells among the myeloblasts. When the degree of differentiation is low (Brugo et al., 1975; Carmichael and Lee, 1977), additional stains, such as Giemsa for eosinophilic granules, PAS for neutrophilic intracytoplasmic granules, and naphthol AS-D chloracetate esterase, may be very helpful (Leder and Stutte, 1975). Reactions for peroxidase and the use of Sudan black B are also recommended. Cytoplasmic basophilia, ascertained with methyl green-pyronine and Giemsa, is lower in immature myeloid cells than it is in immunoblasts.

Immunoblastic ML is not always cytologically monomorphic but may show two main types of
admixture. More commonly, immunoblasts are mixed with a component of plasmablasts and mature plasma cells (Fig. 29). The presence of the latter has been considered a prerequisite for the recognition of a true immunoblastic sarcoma ( Lukes and Collins, 1974 ). On the other hand, this finding may indicate either some trend to a plasmacytic differentiation of the tumour or suggest the origin from a preceding plasmacytic benign or malignant proliferation or from a lymphoplasmacytoid ML. In other instances, immunoblasts are intermingled with malignant centroblasts or even centrocytes, even though a follicular structure is never observed. Macrophages and occasionally epithelioid cells are also found ( Kaiserling, 1977a ), and plentiful reticulum fibre production may be seen. Histiocyte proliferation and fibre production by fibroblastic reticulum cells may possibly be stimulated by the immunoglobulin-producing and immunoglobulin-secreting tumour cells.

Marrow invasion is not uncommon, and immunoblastic leukaemia is characterised by the presence in the blood of large macronucleolated and basophilic immunoblasts ( Fig. 30, inset A ). This leukaemia has been described ( Mathé et al., 1975a ) as type 1 of leukaemic lymphosarcomas. Similarly, previously reported cases of a transformation of ' reticulum cell sarcomas ' into acute leukemia might have been immunoblastic ML ( Lowenbraun et al., 1971 ).

Even though a large proportion of 'histiocytic' and 'undifferentiated pleomorphic' ML retain with marked variability B-lymphocyte markers ( Jaffe et al., 1977 ), it seems understandable that in a number of cases of morphologically characteristic malignant immunoblastic proliferations, immunological markers cannot be visualised and that the cells remain unidentified to a great extent ( Brouet et al., 1975b, 1976; Habeshaw and Stuart, 1975; Jaffe et al., 1977; Kaiserling, 1977a ). In these cases the term immunoblastic ML is not strictly appropriate, and the noncommittal term large lymphoid cell tumour should be applied. It seems, however, that those immunoblastic ML that supervene in patients with previously ascertained B-cell-derived ML may be considered B-cell-derived high-grade malignancies.

Large cell T-type ML ( Jaffe et al., 1975, 1977 ) and specifically T-immunoblastic ML, that is, ML made up of T-cell type transformed lymphocytes ( Habeshaw and Stuart, 1975 ), cannot at present be distinguished with certainty on light microscopy.

Ultrastructurally, neoplastic B-derived immunoblasts ( Fig. 30 ) have a large, round or oval nucleus with dispersed chromatin and one or two prominent nucleoli usually located centrally or occasionally against the nuclear membrane. Rough endoplasmic reticulum is generally scanty, but the cytoplasm is crowded with monoribosomes and many polyribo- somes ( Fig. 30, inset B ). However, some plasmablast-like cells with many rough endoplasmic reticulum cisternae are also present. A well-developed Golgi apparatus and a few mitochondria are found. The diffuse or globular PAS positivity, which is sometimes seen within the cytoplasm on light microscopy, corresponds to immunoglobulin produced by the cells and accumulated in the perinuclear spaces and the cisternae of the rough endoplasmic reticulum.

**Malignant lymphoma, lymphoblastic and acute lymphoblastic leukaemia**

**MALIGNANT LYMPHOMA, LYMPHOBLASTIC CONVOLUTED CELL TYPE**

This entity is characterised by the peculiar morphology especially in sections of a variable proportion of the nuclei of the proliferating atypical cells, which under low power appear round, whereas under high power they disclose a 'cerebriform' appearance due to deep indentations and lobulations almost without intervening cytoplasm ( Fig. 31, inset ) ( Lukes and Collins, 1974; Barcos and Lukes, 1975 ). The percentage of cells of convoluted nuclei varies considerably from case to case and decreases in patients beyond adolescence. In the absence of nuclear convolution the cells are highly immature in appearance and indistinguishable from the lymphoblasts of ALL ( Nathwani et al., 1976 ). This ML used to be classified among the diffuse lymphocytic, poorly differentiated lymphomas, and since it is more frequent in, but not exclusive to, childhood and adolescence ( Rilke et al., 1975 ) it was also called childhood lymphosarcoma ( Sternberg, 1915 ).

In sections of lymph nodes, there is infiltration by a population of cells with a considerable variation in size ( 10-20 μ in diameter ); one of the distinctive features of this ML is the non-cohesiveness of its cells. The nuclear chromatin is delicate, primitive, and 'dusty', and the nucleoli are small and inconspicuous ( Fig. 32 ). The mitotic index is very high, and, in addition, numerous pyknotic nuclei simulate mitotic figures. The cytoplasm is scarce and ill defined. A starry-sky pattern due to actively phagocytosing macrophages may be present. Residual secondary follicles may be found; reticulin fibres are scanty.

In less than half of the cases PAS-positive material is irregularly present in the cytoplasm, usually in the form of circumscribed paranuclear foci or of granules ( Smith et al., 1973; Catovsky et al., 1974b ). In the majority of cases there is a strong positive acid phosphatase reaction in the cells ( Catovsky et al., 1974a; Lennert et al., 1975b, 1975c; Ritter et al., 1975 ), which is restricted to one site ( Fig. 31 ).

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Fig. 30 Malignant lymphoma, immunoblastic. Immunoblasts with oval or round pale nuclei, prominent nucleoli, and scanty rough endoplasmic reticulum. (× 4000) Inset A: ML, immunoblastic; 66-year-old man. Blood smear: WBC 56 × 10^9/l. Circulating immunoblasts 40%. Two immunoblasts. (May Grünwald-Giemsa × 1000). Inset B: Abundant polyribosomes. (× 30,000)

Fig. 31 Malignant lymphoma, lymphoblastic, convoluted cell type. Imprint of cervical lymph node, 30-year-old man. Spot-like acid phosphatase activity (Leder and Stutte, 1975). (× 1000). Inset: circulating 'convoluted' lymphoblast in blood smear. (May Grünwald-Giemsa × 1000)

Fig. 32 Malignant lymphoma, lymphoblastic, convoluted cell type: axillary lymph node, 5-year-old boy. Immature non-cohesive cells with irregularly shaped nuclei, small nucleoli, and scanty cytoplasm. (Giemsa × 1000)
corresponding to the Golgi area and to some adjacent lysosomal granules (Catovsky et al., 1975b). Non-specific esterase, peroxidase, Sudan black B, and alkaline phosphatase are negative.

On electron microscopy the tumour population is formed by cells of different size and nuclear features (Fig. 33). The largest cells have been described (Barcos and Lukes, 1975) as having a nucleus that is more deeply infolded than it is in the smaller ones; however, small cells with convoluted nuclei and large cells with roughly oval non-indentated nuclei are found. In some areas cells with kidney-shaped nuclei are the most frequent. Larger cells generally have more dispersed nuclear chromatin than do the smaller ones. Nucleoli are not prominent. The cytoplasm contains some round mitochondria and a few rough endoplasmic reticulum cisternae. Polyribsomes are abundant, especially in large cells. A well-developed Golgi apparatus (Fig. 33, inset), frequently localised within a nuclear indentation, is observed in a large number of cells. Lysosome-like dense bodies are found, especially in the Golgi region (Fig. 33, inset). Lysosomes and the Golgi apparatus probably correspond to the focal accumulation of acid phosphatase activity observed by light microscopy (Lennert et al., 1975b). The plasma membrane does not present any projection or infolding.

The cell type described, which is characteristic for this ML, is almost identical with that found in cases in which the clinical picture at onset is that of ALL with or without a tumour mass. The latter is quite often represented by a mediastinal tumour in the thymic region, with the same anatomical features described by Sternberg in 1915. Quite often, however, the leukaemic manifestation develops after the appearance of the ML, usually within six months from onset, with high WBC count, at a high incidence of early CNS involvement (Greenberg et al., 1976), and a poor prognosis (Catovsky et al., 1974b; Sen and Borella, 1975; Belpomme et al., 1977), particularly in males (Hann et al., 1977). The distinction between leukaemia and lymphoma, based on the presence of marrow involvement at diagnosis, was not of prognostic significance in one reported series (Coccia et al., 1976).

The cells, whether in the marrow or elsewhere, are of lymphoblasts with E receptors (Borella and Sen, 1973; Smith et al., 1973; Catovsky et al., 1974a; Kaplan et al., 1974; Ritter et al., 1975; Stein et al., 1976) and/or human T-lymphocyte antigen (Kersey et al., 1975; Greaves, 1977). The finding in some cases of complement receptors in addition to the E receptor has been interpreted as an indication that the cells derive from immature T-precursor cells (Stein et al., 1976). The cells do not respond to mitogens.

In one case of T-cell ALL, the suppressor activity was retained by the neoplastic cells (Broder et al., 1978).

The availability of various enzymatic markers (Lancet, 1977), such as terminal deoxynucleotidyl transferase (McCaffrey et al., 1975), combined with immunological differences, are already giving some indication as to the possible heterogeneity of T-cell lymphoblastic leukaemias (Gelfand and Chechik, 1976). On the other hand, it should be pointed out that the clinical evidence of a mediastinal mass and the cytological evidence of a positive acid phosphatase reaction are not specific for a ML and/or leukaemia of the convoluted cell type of thymic origin but may also be observed, although less frequently, in cases of ALL in which the cells do not display immunological markers (Chessells et al., 1977; Seligmann et al., 1977b; Catovsky et al., 1978).

**ACUTE LYMPHOBLASTIC LEUKAEMIA**

The difficulties and contradictions in classifying ALL have been discussed (Flandrin and Bernard, 1975) with special reference to the significance of the cellular size (Binet et al., 1975; Mathé et al., 1976).

In general, the lymphoblasts of ALL are immature lymphoid cells with a high nuclear:cytoplasmic ratio and basophilic cytoplasm without granules. They have been subdivided into two main groups according to their size (Lennert and Mohri, 1971). The small cell type consists of cells that measure about 10 μ in diameter, have slightly irregular nuclei with finely granular chromatin, usually one small nucleolus, and scanty and basophilic cytoplasm. The cells of the large cell type measure about 15 μ in diameter and contain a nucleus with a more regular contour and a prominent nucleolus. Intracytoplasmic coarse diastase-resistant, as well as fine diastase-sensitive PAS-positive granules, were described, and their presence was related to prognosis in some series (Laurie, 1968; Humphrey et al., 1974).

Under the electron microscope there is also a variable morphology (Cawley and Hayhoe, 1973). The nuclear:cytoplasmic ratio is high. The nucleus is seldom kidney-shaped, but mostly it is deeply indented (Fig. 34) and presents nuclear pockets (Fig. 35). Nuclear chromatin is generally dispersed, but there are some chromatin condensations (Fig. 34). However, dense nuclei such as those of small circulating lymphocytes are never observed. Nucleoli are variably developed and occasionally have a prominent nucleolonema. A few peroxidase-negative pleomorphic granules of a lysosomal nature (Fig. 34) are sometimes seen in the cytoplasm. The mitochondria are variable in size, shape, and total number; in some cells they may be abundant. A large accumulation of glycogen (Fig. 34, inset) is
Fig. 33 Malignant lymphoma, lymphoblastic, convoluted cell type. Lymphoblasts with non-idented pale nucleus (A), kidney-shaped nucleus (B), and lobed convoluted nucleus (C). (× 10 000) Inset: Golgi apparatus and lysosome-like dense bodies. (× 30 000)

Fig. 34 Acute lymphoblastic leukaemia. Lymphoblasts with more or less deeply indented nuclei. One cell (A) shows condensed nuclear chromatin, an inconspicuous nucleolus, and a lysosome-like dense granule. (× 12 000) Inset: large accumulation of glycogen. (× 27 000)
characteristic of the leukaemic lymphoblasts (Lennert, 1975), and bundles of microfilaments are sometimes noted. The cytoplasm contains little rough endoplasmic reticulum and is crowded with monoribosomes and some polyribosomes. Differences in ribosomal cytoplasmic content between large and small cell type ALL were reported as the most relevant distinguishing feature; large cells are rich in polyribosomes, whereas in small cells monoribosomes predominate (Kaiserling, 1977a). Membrane-bound dense inclusions of an uncertain nature and autophagic vacuoles are not infrequently observed. A poorly developed Golgi apparatus is present (Fig. 35). Transmission and SEM do not show differences between T and non-T ALL; however, electron microscope cytochemistry may be able to distinguish between null- and T-lymphoblasts. Whereas the strong acid phosphatase reaction of T-ALL blasts is localised within the Golgi apparatus and within the lysosomal granules near the Golgi areas in 30 to 70% of the cells and whereas granules localised in other cytoplasmic areas are generally unreactive, in patients with non-T ALL the reaction is present in one or two granules of less than 10% of the cells, but not within the vesicles of the Golgi apparatus (Catovsky et al., 1975b).

In ALL, the lymph nodes may show a variable degree of distortion of the architecture. While in some cases the enlargement of the lymph node is due to a massive accumulation of cells without marked destruction of the architecture of the lymph node, in other cases the lymph node tissue is entirely replaced by the proliferating cells, as in other diffuse ML (Lennert and Mohri, 1971). The marrow is usually massively infiltrated by tumour cells.

The contribution of morphology, both at the light microscopic cytological and histopathological level as well as at the ultrastructural one, is certainly not satisfactory but immunological and clinical studies seem to be more fruitful. In summary, these investigations have revealed the existence of several types of ALL.

The least common is the monoclonal B-cell leukaemic proliferation of Burkitt lymphoma blasts (less than 3% of all ALL). The report of rare cases of B-cell ALL with the cytological features of poorly differentiated lymphocytic ML (Brouet et al., 1975c) indicates that the patients with ML of germinal centre cell origin (centrocytic?) should be considered at risk for leukaemia and accordingly monitored clinically. Cases of ALL of apparent B-cell origin have been described in adults (Gajl-Peczalska et al., 1974), and rarely, as already mentioned, acute blastic transformation of B-CLL can occur.

The previously described T-ALL seems to be closely linked to the lymphoblastic T-type convoluted cell ML and represents about 25 to 30% of all ALL (Borella and Sen, 1973; Chin et al., 1973; Kersey et al., 1973; Greaves, 1977). For diagnostic purposes E-rosette formation by these lymphoblasts appears to be the most reliable diagnostic marker, while acid phosphatase is somewhat less specific (Chessells et al., 1977), and may be less reliable than the reaction for the intracellular acid α-naphthyl-acetate esterase activity (Müller et al., 1975). It was found that T-cell ALL lymphoblasts present a positivity that is comparable with that of human thymocytes but different from that of T-lymphocytes (Kullenkampff et al., 1977). The considerable difference in prognosis between T-cell ALL and the majority of the other ALL (null-cell) was pointed out by Reid et al. (1977).

The demonstration of the heterogeneity of the group of ALL whose cells do not express any surface characteristics has been given by the identification of a subgroup that reacts with a specific anti-ALL serum (Greaves et al., 1975; Greaves, 1977) and of another that does not (null-cell type). The former is the most frequent type of ALL (65%), and its most important prognostic feature is the level of the initial WBC count (Chessells et al., 1977). Null-cell type ALL still seems to be an inhomogeneous group of diseases.

Histiocytic disorders

A large portion of ML made up of large cells and formerly called reticulum cell sarcomas and termed histiocytic ML in the original Rappaport classification has been shown to derive from various modulation stages of the lymphatic cell. Immunological studies revealed that the majority of large cell ML that originate from a progression of low-grade ML with identifiable markers retain those markers, whereas de novo high-grade ML may have a variety of membrane markers or none at all (Jaffe et al., 1975; Brouet et al., 1976). It was estimated that about 50 to 60% of large cell ML are made up of B-cell, about 10% of T-cell, and more than 30% of null-cell populations (Seligmann et al., 1977a). Thus although the term histiocytic ML is therefore inappropriate in most instances, it could also be that in some cases there is a proliferation of malignant histiocytes (Payne et al., 1977). This could be proved by the demonstration of the functional features of this cell type, such as phagocytosis, enzymatic properties, synthesis of muramidase, and the presence of receptor sites for immunoglobulin and complement on its cell surface (Jaffe et al., 1977; Seligmann et al., 1977a).

Owing to the common feature of having cytoplasmic processes, histiocytes are classified among
Fig. 35  Malignant lymphoma, lymphoblastic. A lymphoblast showing an irregularly lobed nucleus with dispersed chromatin and relatively well-developed nucleolus and Golgi apparatus. (× 13 000)

Fig. 36  Malignant histiocyotosis. An actively phagocytosing histiocyte with large digestive vacuoles. (× 10 000)
reticulum cells. Whereas the histiocytes of germinal centres (‘tingible body macrophages’) and sub sinus phagocytes are phagocytosing reticulum cells, the dendritic, interdigitating, fibroblastic, and dark reticulum cells are not (Kaiserling, 1977a; Stuart, 1975). Further differences between these cell types become evident on electron microscopy and with enzyme histochemistry.

In spite of the anatomical, cytological, functional, and immunological differences between the peripher al lymphatic tissue and the mononuclear phagocyte system and their different derivation (van Furth et al., 1972), it seems appropriate to mention in this context also the neoplastic malignant disorders of the latter, essentially because their true nature can be established, in most instances with reasonable certainty, by morphology alone using light and electron microscopy (Henry, 1975) on lymph node biopsies. A classification of proliferative histiocytic disorders has been proposed on the basis of the unitary concept that all histiocytes or tissue macrophages arise from the BM-derived monocytic series (Cline, 1975). Monoblast-derived macrophages give rise in the tissue to immature ‘A’ macrophages that are capable of cell division and to mature ‘B’ non-replicating macrophages. Neoplastic transformation can affect the normal monocyte-histiocyte line at several levels and produce a spectrum of malignant lesions, which are more or less aggressive according to the degree of differentiation of the proliferating cells. The range of the diseases extends from acute monoblastic leukaemia to malignant histiocytosis, which is considered to represent the proliferation of moderately differentiated cells, and to circumscribed histiocytic proliferations of the most differentiated cells. It has been stressed that the clinical manifestations of the various diseases result from a combined effect of the proliferation of the cells and of their functional manifestations (Cline, 1978). The position among histiocytic proliferative disorders of Hand-Schüller-Christian disease and of eosinophilic granuloma seems debatable, and for them a different derivation has been suggested (Nezelof et al., 1973).

**Malignant Histiocytosis**

This disease is a clinicopathological entity (Byrne and Rappaport, 1973; Berard and Dorfman, 1974; Warnke et al., 1975; Mathé et al., 1976; Rilke et al., 1978a) that is characterised by the malignant progressive and systemic proliferation of histiocytes and of their precursors (Rappaport, 1966). Malignant histiocytosis has been considered the only proved example of a neoplastic disease of ‘true’ histiocytic origin, since it may represent the neoplastic transformation of the histiocytes lining the sinuses of the lymphoreticular system (Berard et al., 1976). It was originally described by Scott and Robb-Smith (1939) as histiocytic medullary reticulosis. The anatomical areas predominantly involved by the disease are the lymph nodes, spleen, liver, marrow, and skin (Byrne and Rappaport, 1973; Warnke et al., 1975). The diagnosis is made by histology on biopsy material but may sometimes be apparent in bone marrow smears.

The most important histological criteria for the diagnosis on lymph nodes, which are the elective site for the identification of the disease, are represented (a) by the presence of a neoplastic proliferation within the peripheral and medullary sinuses and/or within the parenchyma of cells that can be identified as histiocytes because of their cytological features, including phagocytosis, and (b) by the distribution of malignant cells that are predominantly isolated and do not usually form either cords or solid nests. The fibrous capsule of the lymph nodes is not usually infiltrated. The neoplastic histiocytes are not monomorphic because of the mixture in variable proportions of deceptively benign-looking cells with highly atypical ones. The latter have thick nuclear membranes, irregular heterochromatin, and prominent but not very large nucleoli. Phagocytosis is a constant finding, even though quantitatively variable; within the cytoplasm of the histiocytes, erythrocytes, leucocytes, platelets, fat droplets, cellular debris, and PAS-positive granules may be found. Erythropagocytosis in particular may be massive and as such is diagnostic. The number of mitotic figures is variable. Plasma cells and eosinophilic granulocytes are often but not constantly found in a variable amount.

In the spleen, neoplastic proliferation occurs predominantly within the red pulp, while the follicles may persist for a long time and are partially or totally obliterated in the advanced phase of the disease. In the liver, cellular growth is recognisable as an infiltration of the portal spaces and/or of the liver parenchyma with the presence of histiocytes within the liver cords. The finding of histiocytes in marrow is not constant, and invasion may be made up only of small foci of atypical cells. In the skin the atypical histiocytes are more often grouped in the dermis around the skin appendages and the vessels and may also infiltrate the papillary dermis and the subcutaneous tissue.

In addition to the light microscopic findings mentioned, the histiocytic nature of the neoplastic cells is confirmed by enzyme histochemistry, immunology, and electron microscopy. The high cytoplasmic content of acid phosphatase activity (Leder and Stutte, 1975) and of fluoride-resistant non-specific esterase (Li et al., 1972) is easily demonstrated. Immunological markers for monocytes (receptors for
Fig. 37  Malignant histiocytosis. A non-phagocytosing histiocyte with two prominent nucleoli, a large Golgi apparatus, and some primary lysosomes. (× 10 000)

Fig. 38  Malignant histiocytosis. An undifferentiated cell with a horseshoe-shaped nucleus and an inconspicuous nucleolus. Lysosome-like dense bodies are absent. (× 10 000)
the Fc portion of IgG and for complement) have been identified on the cells of malignant histiocytosis (Jaffe et al., 1975).

In our ultrastructural study of six cases of malignant histiocytosis (Lombardi et al., 1978), we observed three types of proliferating cells: actively phagocytosing histiocytes (Fig. 36) with large digestive vacuoles and numerous primary lysosomes; non-phagocytosing histiocytes (Fig. 37) with primary lysosomes only; and undifferentiated cells (Fig. 38) with a few small primary lysosomes. Non-phagocytosing and phagocytosing histiocytes (Figs. 36, 37), which show ultrastructural features similar to those of the A and B macrophages described by Cline (1975), have an oval or deeply indented lobed nucleus with well-developed nucleoli and scarce and margined heterochromatin. Undifferentiated cells (Fig. 38) have a horseshoe-shaped or occasionally a lobed pale nucleus with an inconspicuous nucleolus. All these cell types generally have a scarce, smooth, and rough endoplasmic reticulum and abundant free ribosomes. Some phagocytosing and non-phagocytosing histiocytes with a well-developed rough and smooth endoplasmic reticulum were observed in two cases only. Some multinucleated phagocytosing histiocytes and some cells with features intermediate either between undifferentiated and non-phagocytosing histiocytes or between non-phagocytosing and phagocytosing histiocytes are also observed.

In some cases of circumscribed histiocytic cutaneous and/or nodal neoplasm, similar proliferating cells were observed. Histologically at high magnification the tumour cells could not be distinguished from those found in the sinuses of clear-cut cases of malignant histiocytosis. At low power, however, the higher degree of cohesiveness and the lower degree of atypia represented the main, although qualitatively not well-defined, differences between localised histiocytic tumours and malignant histiocytosis.

Histological differential diagnosis takes into consideration familial haemophagocytic reticulosis (Farquhar et al., 1958), immunoblastic hyperplasia, sinus histiocytosis with massive lymphadenopathy (Rosai and Dorfman, 1969), other histiocytic disorders, high-grade non-Hodgkin's ML, and Hodgkin's disease (Byrne and Rappaport, 1973; Warnke et al., 1975; Rilke et al., 1978a).

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Requests for reprints to: Dr. F. Rilke, Instituto Nazionale per lo Studio e la Cuca dei Tumori, Via Venezia 1, 20133 Milan, Italy.