ON THE PATHOLOGY OF HODGKIN'S DISEASE

by

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ARRANGEMENT OF THE WORK

TEXT OF THESIS VOLUME I

PROTOCOLS AND REFERENCES VOLUME II

PHOTOGRAPHIC ILLUSTRATIONS VOLUME III

DRAWINGS, DIAGRAMS AND GRAPHS VOLUME IV
Hodgkin's disease is the central interest in this study. It is difficult to confine researches strictly to the avowed object because in this disease wider issues obtrude continually. It is common knowledge that Hodgkin's disease affects the reticulo-endothelial system, or more closely, the lymphatic system. These general considerations must be respected and this involves the presentation of summaries of relevant knowledge to form a background to the study of the disease itself.

The work is divided into five main parts. There is first a brief account of the reticulo-endothelial system and its diseases, which is followed by remarks on the lymphatic system. In the third part Hodgkin's disease is reviewed and observations on its morphology and nosological affinities are offered. Next, certain evidences drawn from other diseases are presented in support of arguments and propositions advanced previously, and the final part is reserved for the experimental work on the disease.

There are two appendices of protocols recording the observational work performed. Illustrations supporting the several parts are given, and a final summary concludes the study.
APOLOGIA

'Gutta cavat lapidem non semel sed semper cadendo'

In the warfare that is medicine Hodgkin's disease is an insistent challenge. Figuratively, this uncompromising fortress stands unreduced on the far side of the no-man's-land of pathology, while the redoubts of other diseases succumb to investigation. Those who study Hodgkin's disease will agree that here is no Castra Romana defended by vallum and fossa, but a formidable example of the celebrated third system of Vauban. Every approach is curtained and enfiladed, and the ground is beset with fougasses and chevaux-de-frises. Those who would attack it have chiefly a tradition of failure for their encouragement, yet this should be tempered by reflection on the fates of impregnable fortresses as recent as Festung Europa and Monte Cassino. For most, the assault must be carried out through the smoke of ignorance. It is a recurrent experience that this screen clears as the first bastion is reached, and reveals the objective to have been a mirage, and the raiding party to be still floundering in the fieldworks. This is almost a true similitude but ground is being won, albeit piecemeal. It is a 'salient fact that almost all real knowledge and understanding of this disease has been gained by morbid anatomists who, if the simile be maintained, are the infantry of pathology. The view that properly supported infantry can do anything has long been a classical axiom in military science, and until the equivalents of atomic weapons become available in pathology, morbid anatomy will be the basis of success.

It is submitted that so far as can be foreseen at present final understanding of Hodgkin's disease is more likely to be derived from /the
the consolidation and integration of the morbid anatomists myriad small facts than by any other method.

The present work is primarily an observational study made for this purpose. It is supplemented by information from other diseases and by limited empirical animal experimentation. While it is premature to submit what is properly relegated to the summary, at this stage, the complete failure of the new experimental method tried-out reinforces the decision which was taken.
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PART I

THE RETICULO-ENDOTHELIAL SYSTEM

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THE STRUCTURE OF THE RETICULO-ENDOTHELIAL SYSTEM

Considered collectively as an organ, the reticulo-endothelial system may be described as composed of a stroma and a parenchyma, Piney (1925). This concept is reasonable and helpful, but stereotyped ideas of the stroma and parenchyma relationship in organs like the liver, or salivary glands & etc. must be abandoned for its appreciation. The relation of the neuroglia to nerve cells in the central nervous system presents an approximate homology.

The Stroma

The stromal moiety is fundamentally connective tissue. This may be mesenchyme, general connective tissue, or reticular tissue. Mesenchyme is composed of mutually identical small cells with scanty cytoplasm which are uniformly dispersed. The cells have slender branching processes which are continuous from one cell to another forming a fine open mesh. In adult human material strictly homologous tissue is rare and scanty. A near approach is said to be seen in the nucleus pulposus of the intervertebral discs. These structures, numbering twenty three in all, have received scant attention until comparatively recently but the work of Schmorl confirms the widely quoted report that they are, in part at least, derived from the notochord. Duval, (1900) gave a brief account of their microscopical structure. Collins (1949) described them as consisting of myxoid or chondroid basophile matrix traversed by a few collagenous fibres, and sometimes containing stellate cells like those of a myxoma, or large vesiculated balloon cells resembling those of a chordoma.
In my own studies the nuclei pulposi recovered from elderly people have usually been devoid of cells, but they were demonstrable in children. Physaliphorous change is commonly visible, but tissue morphologically close to mesenchyme is not. A closer approach to embryonic mesenchyme is seen in some myxomata but these are very rare tumours - many supposed examples are merely dropsical degeneration, true mucoid material being absent. It is stated that mesenchymal cells are to be found easily in the vicinity of blood vessels, Robb-Smith (1938). These cells are also described as the adventitial cells of Marchand, resting wandering cells (Maximow), pericytes & etc. I have seldom succeeded in convincing myself of their identity in ordinary routine sections.

The facts are that the methods and material of ordinary human morbid anatomy do not readily yield convincing evidence of persistent mesenchyme in adult tissues. Confirmation of this phenomenon calls for facilities which are not readily attainable, namely absolutely fresh tissue, tissue cultures and much time.

In the adult the mesenchyme is represented by the loose connective tissue and the reticular tissue. The former is composed of fibroblasts or fibrocytes, and histiocytes, with a variable amount of collagen and elastic fibre. Mast cells and pigment-laden cells - probably histiocytes, are sometimes also present. In adipose tissue the fat cells are derived from fibroblastic cells.

The ubiquitous general connective tissue of the body can be regarded as relatively generalised and undifferentiated tissue, preserving in some degree an embryonic potentiality for undergoing differentiation into more
highly organised supporting structures, generally by some form of fibre production. To this extent it has been regarded as a persistence of the embryonic mesoderm (mesenchyme).

The reticular tissue is nearer to the parent mesenchyme. The same general arrangement obtains but the cells are larger and the mesh shows local differences. Although the tissue is described as net-like it is more accurately represented by a sponge which is the skeletal remnant of a porifer, \textit{(genus Euspongia)}. The mesh is formed of protoplasm and a variable amount of fine argyrophile fibre called reticulin. This fibre material is often manifestly in direct continuation with collagen but unlike the latter its fibres run singly, anastomose freely, and are of uneven thickness. Triturated collagen is argyrophile, its component fibrillae take on the silver. It is unlikely that reticulin is a specific substance, and the argyrophile property is certainly not an exclusive possession of these fibres. Elastic fibres (elastin) are not present in reticular tissue itself. In some texts the term retiform tissue is used and in others the words reticulum or pulp are employed as equivalents. The last named is apt to lead to confusion.

In paraffin sections stained with haematoxylin and eosin it is usually difficult to identify this ground tissue in normal lymph nodes, spleen, bone marrow & etc. Silver impregnation is often difficult to interpret. This can be helped by substituting the orthodox counterstains saffranin or neutral red, with haematoxylin of light intensity. Another device is to mount two adjacent sections on the same slide, one being impregnated with silver, and the other stained with haematoxylin and /eosin
Comparison is thus easily made. It must be emphasized that the silver impregnation is only a partial picture since the ground protoplasm is not stained. The most convincing demonstration of the protoplasmic continuity of reticular tissue is obtained by the old device of 'pencilling', Renvier (1889); Jolly (1922). These preparations are easily seen unstained, but light aniline stains may be used.

The proportion of stromal tissue varies in different sites, it is relatively high in the spleen and lymphoid tissue and low in the bone marrow and liver.

The component 'cells' of the reticular tissue, which is properly a plasmodium, are usually termed reticulum cells. There are many other names as will be seen later. There is certainly confused thinking concerning the reticulum cell. It is obvious that many authors refer to reticulum cells when they actually mean reticulum cell nuclei. This error is understandable because the cytoplasm is generally so inconspicuous and faint-staining. The usage is common and is apparently condoned. The characterisation of this cell is difficult because although quite precise definition is available from many sources the title has a certain omnibus propensity. The term, actually as 'reticular cell', was first introduced by Ribbert (1889). He applied it to the true reticulum cells of lymphoid tissue to distinguish them from the modified reticulum cells of the lymph sinuses which were formerly more definitely termed endothelial cells. It is convenient to allow a degree of elasticity in the descriptive morphology of this cell, otherwise an host of pro- and meta- forms etc. and many named variants

/complicate
complicate the issue. Thus, in lymphoid tissue the 'cells' of the pale centres of germinal follicles, the reticular cells of Ribbert, the proliferating cells of Marchand, the 'endothelial' or 'littoral' cells of the sinuses are all reticulum cells, despite their individual differences. Similarly the various primitive blood cells, e.g. haemohistioblast, haemocytoblast, common lymphoid stem cell, lymphoidocyte & etc. are all considered as reticulum cells. This wide application is open to criticisms but if the varieties of reticulum are imagined as siblings of a family - mutually similar and yet dissimilar at the same time, the paradox is acceptable.

The Parenchyma

This part of the complex rather defies attempts to consider it as a whole because the individual scions of it subserve very different functions. There are however several features which encourage some reflection.

This component of the system is entirely cellular. Formed elements of cells, at least in the generally understood sense, do not contribute. The minor exceptions alluded to are the surface films which may be shed from macrophages, lymphocytes, or plasma cells, in immunity development. The cells of the reticulo-endothelial parenchyma are less mutually coherent than any others in the body, many indeed are entirely free. This characteristic is interesting because it is balanced by a functional individuality, the morphological and functional versatility of the cells in this system far surpassing that observed in any other tissue. The majority of the free forms have independent mobility, an archaic property which betokens much; even the relatively fixed cells /have
have potential to move provided an appropriate stimulus is offered.

The parenchyma is conveniently described for the purposes here under four main heads. These are the erythron (Boycott), the leukon, the macrophageion, and the lymphocytogeion. (Approbation of these last three neologisms is not anticipated but they avoid unwieldy circumlocutions).

There is general agreement that series of cells can be recognised which ultimately form erythrocytes and myeloid series leucocytes. Specialisation in function and morphology is carried to the extreme in the former and it seems certain that a stage of differentiation is soon attained in the progenitors whereby erythrocytes only can be produced thereafter. The myeloid leucocyte is similar but specialisation is not carried nearly so far. The end-cell still retains its power of independent motion, of phagocytosis and is nucleated.

The macrophageion is nearest of all to its fundamental prototype. The parent tissue is not defined so closely as that of the former two, neither in topography nor structure. It is further complicated by the undoubted contribution of functionally identical cells from non-mesodermal sources.

Finally there is the lymphocytogeion. This is the least understood of all. It will be discussed more fully later but the extraordinary mystery of lymphoid tissue which collectively would yield a mass approaching the liver in size (Drinker), and which is so deceptively inert in appearance is a most baffling enigma.

/Indeterminate
Indeterminate cells also contribute to the parenchyma. To observers who are ordinary mortals, their proportion is usually uncomfortably high. Most are doubtless transition forms in exasperating disguises but others may be visitors, child-cells lost in the crowd, or captives.

The mutual relationships of the cells in this system are difficult to study. The problems are inherently very complex and are not amenable to easy investigation. A very confusing nomenclature has developed which makes the consolidation and integration of individual researches most difficult. It is important to appreciate that with the exception of blood-formation, little has been contributed on these problems in the human system. Also, by virtue of the paradox that the anatomists with the deepest experience of human tissues are morbid anatomists, more is understood concerning some aspects of the system in disease than in health. The identity and nomenclature of many of the immature cells of the system are not yet satisfactorily agreed. The mutual identities of cells which might be regarded as equivalent in different species of animals are also far from established. The subject has prompted a great amount of work much of which is highly controversial. Adding to these difficulties inconsistency pervades even the best writings. These soon impress the tyro who makes an honest and critical approach to the subject, and unless he is exceptionally fortunate he also discovers that very few can help him.

Certain peculiar features attend the origin and cellular maintenance of the reticulo-endothelial system. It is axiomatic that each human individual
individual starts existence as the fusion product of two haploid cells and therefore every cell in the body must own to this common ancestor. Specific differentiation of almost all descendants of this first cell follows with eventual production of all the different cells of the adult. This differentiation is accomplished remarkably early and thereafter the increase is chiefly on the homoplastic plan; that is to say muscle increases by proliferation of pre-existing muscle cells & etc. In many tissues these fully differentiated cells are end-products; the nerve cells of the central nervous system enjoy an impregnable reputation for immutability, even to the absence of propagation.

The connective tissue from which the reticulo-endothelial system springs deviates from this general plan, but the difference is not so much one of kind as one of degree. There is good evidence that environment and functional demands determine the individual morphological varieties of cell produced, the genetic influence being less concerned with the ultimate form of the cells than with endowing them with ability to react to a wide functional spectrum.

It is a general rule that cells which differentiate to subserve a particular function lose some of their ability to de-differentiate or metamorphose, and this loss is increased as that function becomes more complex or specialised. This law is evaded by the reticulo-endothelial system cells on a remarkable scale; the erythrocytes and polymorphonuclear leucocytes are probably the only two real end-cells of the entire system. Neither is capable of cell-division and so far as the body economy is concerned they are expendable. The remainder, even the...
mature fibrocyte and the ever-incongruous lymphocyte can probably adopt the habits of most other cells of the macrophage system. This lability must be regarded as a system characteristic. One important outcome of these curious properties is that in tissue of this system a departure is made from the stroma and parenchyma relationship observed in ordinary organs. In the reticulo-endothelial system the parenchyma arises from the stroma. This is a fundamental peculiarity, and remains a central fact even if the issue is complicated by the contentions of the polyphyletic or monophyletic schools. These latter arguments have acquired undue advertisement; what they chiefly seek to determine is whether there are maturation levels in the earliest precursor cells after which homoplastic generation of one type of cell is finally determined. Good evidence exists in favour of both theories. The inflammatory cellular response, the totality of aplastic anaemia, and the phenomenon of myeloid metaplasia favour the monophyletic view; the extreme rarity of mixed leukaemias, the polyphyletic one. It is reasonable to accept both as correct, they are in reality far less irreconcilable than is often supposed. One can merely presume that under appropriate circumstances - of which knowledge is very restricted - one or other path is taken. Comprehensive studies on the genetic relationships of the reticulo-endothelial cells have been made. Among the most valuable are those of Maximow and his colleagues. Much of their work has consolidated and integrated in Maximow' well-known review in 1924. It is to be noted however that these findings were almost exclusively derived from fresh material, a method which has very restricted application to man. It is
apparent that in some quarters uncritical transference of these results to man has been made with the result that generalisations have been formulated without a really satisfactory basis.

The recognition of the fundamental cells of the system in man presents much difficulty to the beginner because the discrepancies between authoritative descriptions are serious, e.g. Robb-Smith (1938), and Hadfield and Garrod (1943), describe undifferentiated mesenchymal cells and reticulum cells so differently that their mutual identities are quite irreconcilable. Many schemata describing the inter-relationships of the cells of the system have been prepared, e.g. Maximow (1924), Hadfield and Garrod (1943), and Gillman, Gillman, and Gilbert (1949), but it is difficult to assess how much can be applied to man. The metamorphoses believed to occur in the human reticulo-endothelial system which are of particular interest in the study of lymphoid tissue are summarized in the following table.
<table>
<thead>
<tr>
<th>Precursor cell.</th>
<th>Differentiation product.</th>
<th>Authority.</th>
</tr>
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<tbody>
<tr>
<td>Reticulum cell.</td>
<td>Fibroblast, fibrocyte</td>
<td>General.</td>
</tr>
<tr>
<td></td>
<td>Epithelioid (or endothelioid) cell.</td>
<td>&quot;</td>
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<tr>
<td></td>
<td>Littoral cell. (Lymph sinus).</td>
<td>&quot;</td>
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<tr>
<td></td>
<td>Macrophage.</td>
<td>&quot;</td>
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<td></td>
<td>Monocyte (histiocyte).</td>
<td>&quot;</td>
</tr>
<tr>
<td></td>
<td>Lymphocyte series.</td>
<td>&quot;</td>
</tr>
<tr>
<td>Lymphocyte.</td>
<td>Plasma cell (Marshalko type).</td>
<td>&quot;</td>
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<tr>
<td></td>
<td>Eosinophil.</td>
<td>Dominici; Pullinger.</td>
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<tr>
<td></td>
<td>Blood vascular endothelium.</td>
<td>Moschowitz.</td>
</tr>
<tr>
<td></td>
<td>Macrophage.</td>
<td>Downey and Stasney.</td>
</tr>
<tr>
<td></td>
<td>Erythron.</td>
<td>Jordan; Yoffee.</td>
</tr>
<tr>
<td>Macrophages.</td>
<td>Fibroblast, fibrocyte.</td>
<td>General.</td>
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</table>

This table outlines some of the simpler relationships for which evidence has been found; many other genealogies are derived, but are less well endorsed for the human species.
INTRODUCTORY REMARKS ON THE DISEASES OF THE
RETICULO-ENDOTHELIAL SYSTEM

It is established practice in Special Pathology to study the
diseases of the individual systems separately. The method is generally
satisfactory, even if it sometimes loses its applicability because of its exclusiveness. In practice it serves well and is much easier to
apply than the older and more precise system of the textures.

On general grounds the application of the principle to the reticulo-
endothelial system is quite orthodox. The system now emancipated from
its restrictive functional definition, has, despite its anatomical
diffuseness, a defined topography.

In systemic diseases it is common to find a degree of localisation
of the morbid process. Indeed it happens occasionally that the lesion
is solitary and confined almost rigidly to one site. This extreme state
of affairs is illustrated readily in examples of peptic ulceration of the
alimentary tract. Here it is common to find a single lesion, and the
remainder of the system often natural.

On other occasions while a disease apparently affects one part
predominantly it is in reality much more generalised. Formerly endo-
carditis was stressed as the all-important lesion of rheumatic diseases
of the circulatory system, valvular defects resulting therefrom were
considered the most significant results. Later the discovery by Aschoff
that the connective tissue interstitium of the heart was affected,
together with the clinical insistence on gross myocardial dysfunction
realised the concept of pancarditis. This in turn has given way to the
idea that the disease is primarily a general connective tissue or even a
diffuse collagen affection; a belief supported by the results of
careful scrutiny which disclosed blood vessel and other inconspicuous
lesions which had been previously overlooked. On the other hand systemic
disease may be truly generalised. A familiar example is arteriosclerosis
of the ordinary type which invariably affects all the arterial vessels.
In like manner several of the primary muscular dystrophies involve all
the muscles in some degree.

Analogous variations of involvement are exhibited in disease of the
reticulo-endothelial system but in general, systemic participation in
the morbid process is regarded as the rule. Numerically the vast
majority of alterations in the reticulo-endothelial system are secondary
in character; hardly deserving the appellation disease but rather
falling into the category of reactive or adaptive change. The
universality of participation of this widely scattered system in these
reactive states cannot usually be assessed, in man at any rate, with
much accuracy. This is particularly true when the causes evoking the
reaction are transient or trivial. Pronouncements such as 'general
reticulo-endothelial hyperplasia' while reasonable on theoretical grounds
are in actual practice little more than a shrewd guess. Even where the
stimulant factor is known or presumed on good grounds, and is operative
over a protracted period the interpretation of the results in the
system is not easy. Thus in the peculiar metabolic maladies, the
lipoidoses; in chronic haemolytic anaemia; in pernicious anaemia; in
haemochromatosis & etc., considerable individual variation of the
reaction pattern is well attested fact. These anomalies have not received the attention they deserve, they are probably of considerable significance. The individual response of different people's macrophage systems is as bewildering as the apparently fortuitous distribution of metastases in individual instances of malignant disease.

The primary diseases of the system are the chief concern in this work. Here they may be defined as non-altruistic proliferative conditions arising in it which are not obviously brought about by disease in any other system, and for which it is not possible to assign any specific cause. This category will manifestly include true tumours but there are also conditions which at present do not enjoy universal recognition as neoplasm despite their tumour-like character. These are usually termed primary reticuloses. There is an impression prevalent that the primary diseases of the system have a basic common morbid anatomy related to the sites of location of immature mesenchyme. Thus enlargement of lymph nodes, enlargement of the spleen, involvement of bone marrow and enlargement of the liver are frequently all present. As a working generalisation this is probably fairly true but at the same time it is an overworked hypothesis. Topical variation of involvement is quite evident in the primary maladies of the reticulo-endothelial system just as in other systems.

To the pathologist the problem of greatest interest presented by primary disease of the reticulo-endothelial system is unquestionably its status. This question is complex and very difficult, and it is pertinent at this juncture to discuss the meaning of tumour, and to
present some general points concerning tumour growth in this system.

Tumour, in its proper pathological sense is very difficult to define. Nicholson (1925) maintained, 'That it is impossible to define a tumour. Whenever we look we see that tumours exhibit no differences in kind but only differences in degree – and these often of the slightest – from the other tissues of the body. I have tried for years to formulate a definition, but have failed. Others have been bolder. I shall not weary you with their definitions every one of which breaks down at one or more points'. This is a discouraging reflection but at the same time it overlooks the limitations inherent in definition, it is hard to believe that all experience of an idea or thing can ever be accurately expressed in a few neat words. In any event it would tax even a mind like Nicholson's to define what constitute differences of kind and differences of degree; the nuance can be very delicate. If this limitation is acknowledged then Nicholson is correct and it would be futile to attempt definition; on the other hand it is possible by collating a descriptive account of observed facts to render an imperfect definition which has practical value. This in turn may be expressed in quintessence avoiding at least the grosser errors.

Commencing with Cohnheim a long series of definitions have been submitted and it is interesting to examine some representative examples. Cohnheim maintained that a tumour was 'a circumscribed atypical production of tissue from a matrix of superabundant or erratic deposit of embryonic elements'. This became obsolete when the 'cell-nest' theory was proved inapplicable to tumours in general. Further it is doubtful whether
whether circumscription can be taken to mean more than apartness. Lubarsch included 'all spontaneously arising tissue growths which while typical in form differ in histological structure from the originating tissue, and, in spite of organic connection with the body, pursue an autonomous course, rarely to the benefit of the whole body'. As far as the practical morbid anatomist is concerned this is a good definition but with the advent of newer knowledge it is losing its value. (Still, few experimental pathologists would be ready to assign specific causes in individual cases of human cancer). It is unlikely that Lubarsch referred only to histiomata in his usage 'tissue growths'.

Ribbert regarded tumours as 'self-contained, dependent upon the organism for their nourishment but otherwise largely if not quite independant; corresponding more or less but never absolutely with the tissues of the natural body and presenting no typical limit of growth'. Serious doubt is now cast on whether tumours are self-contained; their non-correspondence with natural tissues may be theoretically true but sometimes their morphological identity can be treacherous. The paradoxical term benign metastasizing goitre is not retained for nothing, and granulation tissue in bone is only separable from sarcoma with assurance - by the really clever - or the foolish. The phrase 'no typical limit of growth' is extremely neat because it allows for the spontaneous cessation which is sometimes apparent. Birch-Hirschfields' definition is similar to Ribbert's but introduces the idea of progressive new growth. Ziegler submitted that 'a tumour is a new formation of
tissue possessing an atypical structure; not exercising any function of service to the body and presenting no typical limit of growth'. This approaches closer to the modern definitions.

Powell-White defined a tumour as follows - 'A tumour proper is a mass of cell tissues or organs resembling those normally present in the body but arranged atypically. It grows at the expense of the body without at the same time subserving any useful function'. This fails to distinguish tumours from malformations (Willis, 1948). But this objection assumes that the latter are distinguishable with certainty, which it is respectfully submitted is sometimes a moot point. The limited capacity for aberrant growth displayed by Albrecht's hamartoma could be regarded as a type of link between the two.

Prudden, Ewing, and Muir concur in the definition of tumour as being 'any autonomous new growth'. This brevity has been condemned, yet autonymity does seem the only feature common to all neoplasms. It has the advantage of course that the less one says, then the less one can be held to, and evades the issue of defining autonymity. This autonymity is very complex, it means that the new growth is a law unto itself in innumerable ways. To interpret it as meaning that tumour disregards the 'laws of normal growth' - which are neither formulated nor understood - is a serious understatement.

Willis submits the most recent authoritative definition namely, 'A tumour is an abnormal mass of tissue; the growth of which exceeds and is uncoordinated with that of the normal tissues and persists in the same excessive manner after cessation of the stimuli which evoked the change'.

/Most
Most emphasis is laid on this last feature which Willis terms the irreversible neoplastic habit of growth. This is in accordance with the facts of experimental carcinogenesis and the author leads good evidence that the time lag in human cancer development after exposure to known carcinogenic hazards is dependant on the same principle. On the other hand the aetiology of most human cancers is still guess-work.

It is important to acknowledge the extraordinary amount of experience, learning, and ingenuity displayed by the originators of these definitions. With a few exceptions all of these statements of what constitutes tumour are basically the same but different facets of the peculiarity of neoplasm are emphasized.

It is invidious to be forced to choose and then permanently adopt any particular one. My interest in tumours is primarily concerned with the ability to recognise them and while I admit much attraction towards the old definitions especially those of Lubarsch and Ziegler, I believe that Muir's 'any autonomous new growth' is the safest. Peculiar new growth is the central idea. It can be applied widely - to tissue, cells, or free cells displaying the change, and displaying it in any degree.

Finally, there is a useful and indeed much used term - 'tumour-like'. So far as I am aware no one attempts to define this beyond calling it blastomatoid, which is like calling the Earth a Geoid. We all labour under the judgment of Ruskin - that it is the hardest thing in the world to put into words exactly what you mean. This, with a sombre background of ignorance, inexperience, and imperfect observation, makes the task even harder. The eye and the brain can recognise appearances accurately
without us being actively conscious of the processes involved. A normally intelligent and interested child can be taught to recognise the mushroom (Agaricus campestris) with infallibility in a matter of a few days; and the only reasons he can offer for his identification is that it is a mushroom. Analogous situations, on a more sophisticated level, apply to the morbid anatomist. 'Yes, it is tumour'. To which the cynical colleague says 'Well, would you stand up in a court of law and defend that statement?' At this point frailty and abject inability to give reasons which sound convincing begin their pernicious work, and a compromise, 'tumour-like' may be substituted.

This is not to be condemned too harshly, provided it is not simply used as an easy way out of every difficulty. The diagnosis of tumour, malignant tumour, is far too serious a matter for doubt. The criteria must be met properly every time. Even so - disease does not always play fair. The gist of the matter is that microscopically tumours nearly all have a 'look' about them which is peculiarly difficult to describe but which the initiated appreciate.

There are several aspects of tumour growth which are particularly worthy of reflection in relation to tumours of the reticulo-endothelial apparatus.

As Willis (1948) has pointed out, Cohnheim's cell-nest hypothesis, despite its abandonment, has left two persistent corollaries. The first is the belief in unicentric origin. For a long period this was accepted as axiomatic and apparently with good reason, the majority of tumours in many organs and tissues are single. On the other hand many...
typically multiple tumours exist, for example duct papillomata of the breast, cancers arising in polyposis coli, and myomata of the uterus.

A further evidence of fields of origin is afforded by recurrence of tumour growth when microscopical examination proves the initial resection to have been adequate. It is suggested that fields of origin can explain autochthonous growth. This phenomenon is highly characteristic of tumours of the reticulo-endothelial system. If the lymphoid tissue collectively is envisaged as an organ which is virtually a single mass of identical tissue it is not difficult to rationalise the idea.

The second corollary is that tumours are purely intrinsic. In human pathology it is certainly rare to encounter sarcomatous change in the stroma of epithelial tumours but lesser degrees of 'taking up' of normal tissue are very probable, and, as Willis maintains, such a phenomenon is explicable on the potential present in a field of origin. That this occurs in lymphoid tissue is certain. It is unorthodox to recognise local Hodgkin's disease in part of a lymph node but it does occur. An explanation for these two facts lies in this hypothesis.

Metastases are common in human malignant neoplastic disease. There is considerable understanding of how they occur but very little interest in the reason for them. It is moderately true to regard tumours as parasites. These sycophantic forms of life will take the most fantastically prodigal steps to ensure their own survival. Possibly lymphoid tissue tumours are more cunning than the others — they do not need to metastasize, they rely on autochthonous growths to further their distribution. (On occasion the whim to metastasize by orthodox /mechanisms
mechanisms may be exercised).

One of the commonest forms of tumour is the spheroidal nodule and it is hoped to shew that this primitive formation is common in many types of lymphoid tissue tumour even if ephemeral and elusive.

There are two further points of general interest. The first is that the process of mitosis is relatively fast, it is estimated that it occupies 1-2 hours in man. Accordingly even one mitotic figure in every high power field reflects considerable growth activity. This proportion is about the minimum observed in many reticulo-endothelial system tumours.

The other point is that it is doubtful whether cells in mitosis can be recognised with certainty. It is perhaps a naïve reminder but they often look singularly alike. Ellermann (1923) led evidence that specific mitotic angles characterised each type of cell, this criterion is however very hard to apply.

The cause of death in neoplastic disease is sometimes obvious enough, e.g. haemorrhage, inflammation, starvation, suffocation, & etc. Often it is not. It is suggested that in lymphoid tissue tumour, death is often the result of a virtual ablation of the lymphoid tissue. The normal lymphoid tissue is reduced to a level incompatible with the maintenance of life, by either anatomical or functional destruction.
PRIMARY RETICULOSIS

The notion that a morbid process might exist intermediate between tumour and inflammation was first advanced by Craigie in 1828. This interjection appeared in his considerations on 'Vascular sarcoma'; the description of the hard variety of this condition has been interpreted by Fox (1926), and others as possibly a reference to Hodgkin's disease. The account however was anecdotal. In the second edition of his 'Elements' (1848), significant new matter was not added concerning this entity, nor was the idea elaborated. This writer has a further claim to priority in this subject which is little known. In 1845 he contributed a paper on two cases of 'Puriform absorption from the Spleen'. He gave a good account of the clinical course and morbid anatomy of two cases which were undoubtedly examples of leukaemia. He noted that in the terminal phase of the illness a severe exacerbation developed and he attributed this to a suppurative episode. Arguing from this, and the fact that abscess of the spleen is of great rarity, he expressed the belief that the 'puriform blood', which he had confirmed by microscopy, originated in that organ and emanated therefrom instead of remaining local. It is remarkable how closely this result, furnished by reasoning, anticipated the true state of affairs. The second of Craigie's patients, John Menteith, formed the basis of a paper published synchronously in the same Journal by John Hughes Bennet, whom MacCallum (1928) quoted as the discoverer of leukaemia. Bennet described the instance as one of 'Suppuration in the blood'. In this he reverted to the old idea of Piorry of an haematitis and clearly supported this curious hypothesis. Later in the same year

/Virchow
Virchow described similar cases and recognised the lymphoid nature of the cells in one type and their granular character in the other. In addition he separated the general syndrome into leukaemic and aleukaemic groups. Later, in 1865, Cohnheim described the latter as pseudo-leukamie. It is interesting that Virchow considered that leukaemia was an expression of a perverted erythropoiesis, whereby the invariably attendant anaemia was neatly explained, a view tenable only so long as this metamorphosis passed as a physiological postulate. This erroneous premiss was abrogated by Neumann, (1879); particular credit for Neumann's recognition of the bone marrow hyperplasia in myeloid leukaemia was awarded by Cohnheim in the second edition of his lectures.

By this time, 1882, Cohnheim was manifestly advocating recognition of systematized hyperplasias of the haematopoietic tissues as the underlying cause of this group of diseases. Moreover it seems very probable that his invention of 'pseudo-leukamie' reflected the application of this idea. The equivalent French terms, 'lymphocytémies frustes', 'formes frustes' Jolly, (1898) convey the same sense. While most cases referred to under this head were probably aleukaemic lymphatic leukaemia, Hodgkin's disease was specifically included by Cohnheim himself. In 1900 Dominici put forward the brilliant suggestion that the 'metastases' in these conditions were due to the local growth of cells at these sites; these cells having powers of differentiation which were evoked in the disease. After this point was gained at a time when much authoritative belief (Ehrlich et alii) favoured the idea that leukaemia and allied disorders were orthodox neoplasms; the morbid process reticulosis appears to have
attracted recognition but doubtful acceptance. The French certainly appear to have had a shrewd percipience of these peculiar aleukaemic hyperplasias; records preserved from this period at certain Paris hospitals include provisional classifications closely similar to those advocated much later in accessible literature. This is a sidelight on the parochial proclivity of the Catholic Gaul! Kastiras, (1945).

The term reticulosis requires elucidation. This form, reticulose or reticulosis, was introduced by Letterer in 1924. Etymologically it is unsatisfactory, Willis, (1948), but is taken to mean proliferation of reticulum. Other forms, some with restricted meanings e.g. leukosis, myelosis, lymphadenosis, reticulo-endotheliosis etc. have not come into wide use.

The following abstracts from British and Empire literature are quoted to indicate the progressive attempts to formulate a definition of reticulosis.

MacCallum, (1928), did not make use of the term but employed the older 'aleukaemic lymphadenosis', and throughout he used the word hyperplasia unqualified.

Pullinger, (1932), did not actually define reticulosis but suggested that 'a group of disease of the reticulum existed in which proliferation was possible into one or several of the possible progeny'.

Rose, (1933), did not define the condition, but, following MacCallum, referred to 'the systemic hyperplasias of the blood-forming organs'.

/Robb-Smith
Robb-Smith, (1933), submitted the following definition.

'A progressive hyperplasia of reticular tissue with differentiation to one or more cell types. It is commonly systematized, that is to say it affects throughout the body tissue homologous to that affected in the lymph node. Further free cellular elements may be present in the circulating blood (leukaemia). Owing to the proliferation arising at many points there may be apparent infiltration but stromal destruction is not seen and dissemination by embolism (proved by emboli in vessels) is rare'.

It is seen at once that definition is difficult and even in the full exposition of Robb-Smith there is still doubt about the specificity of the process.

The morbid process is unique, it occurs solely in tissue of the reticulo-endothelial system. Disease processes restricted to one system are rare in general pathology, the only common example being dental caries, to which there seems no analogous change elsewhere. The rare primary myopathies, in so far as they have been studied and understood, furnish another possible exception. Mental disease may represent isolated systemic disease, but it is too abstract to discuss in the present state of knowledge. The old tetrad; malformation, trauma, inflammation and tumour cover most pathological contingencies in all the systems. The invocation of a new special morbid process is therefore hazardous. The reticulo-endothelial system is certainly unique, but it does not follow that it has pathological changes of its own. The ordinary pathological changes may be merely rendered unfamiliar by reason
of the peculiarity of the system.

It is equivocal what is meant by progressive hyperplasia. The word hyperplasia was invented by Virchow to mean an increase in the number of the cells of a part in response to demand. In ordinary usage this is tacitly accepted as physiological and purposeful or at least altruistic. Further it seems generally understood to connote homoplastic generation. It’s quite properly applied to haemopoietic tissues, e.g. the increase of red bone marrow after haemorrhage; and so must cover the maturation aspect of this process as well. In the definition it is not clear whether progressive means perpetual or autonomous. There is not any indication whether it differs from the continual new blood formation which is maintained throughout life or if it means that the process, once initiated is self-maintaining (i.e. without continued action of the genetic stimulus). If the latter, it is reasonable to say it is tumour growth. Most protagonists of reticulosis avoid close discussion about this kind of hyperplasia; indeed certain writers, e.g. Gorer, (1946), avoid the term hyperplasia, substituting the safer word 'proliferation'.

It is in the character of systematisation that one of the most difficult inconsistencies in the idea of reticulosis lies. Now while it is admitted that in Robb-Smith's definition that this is to be expected only commonly - not invariably, it is understood by many protagonists as an inherent feature. With a few exceptions, the lesions in this group of diseases do not readily affect the general connective tissue of the body where reticular tissue abounds, nor are they to be found in those /inconspicuous
inconspicuous deposits of persistent early mesenchyme which the adult body still harbours. My chief interest has been concentrated on Hodgkin's disease. In my necropsy series of cases of this disease, the omentum, which is a rich, indeed a classical source of reticular tissue, was never affected except in definite lymphoid tissue therein. I examined the nuclei pulposi of the intervertebral discs on a limited scale in the series and never found deposits of the new tissue in them. Fascial planes were equally negative. The rarity of skin lesions is well accredited, and yet collectively the integument contains an enormous number of reticulo-endothelial cells. Goldman and Victor (1945) recorded skin nodules in only two cases out of a total of 319, (0.62 percent). Further observations on the location of the lesions will be offered later but the view that real systematisation occurs is not impressively illustrated in the majority of reticuloses.

The meaning of apparent infiltration is dubious and rests chiefly on personal interpretation. There is, in certain cases, very real difficulty in deciding whether true infiltration has occurred; but it is equally difficult to prove that it has not. In ordinary tumours infiltration is one of the most significant microscopical criteria of malignancy. Here the ability of all leucocytes to move individually complicates interpretation. Likewise their undoubted local development in ectopic sites may invalidate an opinion that they are truly metastatic.

It has long been assumed that the entrance of tumour cells into the lumina of vascular channels is a natural and exclusive consequence of their vital property of invasive growth.
A corollary of practical value is that the finding of tumour cells in blood or lymph vessels reinforces the diagnosis of malignancy. However embolism of normal tissue cells is well known, (Young and Griffiths, 1950). Familiar examples are seen where trophoblastic elements are caught up in the lungs, and where liver cells are seen lying free in the hepatic vein radicles.

This reduces the absolute value of the finding in tumour diagnosis.

It is still true however that blood-borne dissemination is a very important feature of many, if not most, malignant tumours, whatever mechanism precipitates entry. Thus an admission that reticuloses may spread by such a route is suspect.

The relationship with acknowledged tumour is very close. It is certain that individual examples of reticulosis undergo change into frank malignant tumour. This has been recorded in many cases by the study of serial biopsies and subsequent necropsy. At what point is one to draw a line between the two processes? — or, is there a line at all? It might reasonably be held that all tumours are malignant by virtue of their being autonomous new growths but the grade of malignancy can range from exceedingly low to very high. This is justifiable because from a simple point of view all tumours compete with the host's tissues, some in the mildest fashion possible, others with aggressive avidity.

The status of certain reticuloses stands in acknowledged doubt. The disease described first by Brill, Baehr, and Rosenthal in 1925 is a good example. This initially bore the non-committal title 'splenomegalia lymphatica hyperplastica'. (Some pathologists refer to it as Brill's disease;
disease; in so far as eponymic titles are an excellent idea this is good, but confusion may arise with the Rickettsial disease of the same name, cf. the medley of von Recklinghausen's diseases). At present in Great Britain it is better known as lymphoid follicular reticulosis or giant follicular reticulosis, but in the U.S.A. the terms follicular lymphoma or follicular lymphoblastoma are used instead. More remains to be said concerning Hodgkin's disease, but only a few enthusiasts really call it fibro-myeloid medullary reticulosis and very few fibro-myeloid-histiocytic medullary reticulosis. Mallory's title for it (1913), scirrhous lymphoblastoma, is also little used, not because the tumour theory is unsupported but for the good reason that convention has narrowed down the import of lymphoblast. Lymphogranuloma is however still widely used.

In many respects the leukaemias conform better than most other reticuloses to the postulates in the definition. The peculiarities of these diseases are conducive to this. Chronic leukaemia is much commoner than the acute variety. In the former the majority of the leucocytes, whether granular or lymphocytic are usually very close to normal cells in morphology. Hyperplasia is therefore used with much justification to account for their supply. In chronic myeloid leukaemia it is exceptionally rare to encounter localised tumour (chloroma) and the gross generalised granulopoiesis is definite enough in most cases. On the other hand the inherent migratory properties of these leucocytes may conceivably account for this absence of localised tumours.

In chronic lymphatic leukaemia the apparent identity of the
lymphocytes with normal ones is even closer. Here the generalised lymphadenopathy and splenomegaly could be reasonably interpreted as the result of hyperplasia. But, if a section of one of these lymph nodes is examined one pathologist will report it as medullary reticulosis of lymphoid type and another as lymphosarcoma. There is not any reliable criterion offered to tell the two apart; it is open to question whether these are not simply two histological names for the same thing.

In the acute leukaemias the supporters of reticulosis do not press their claims; the balance is too heavily weighted in favour of tumour. In the myeloid form localised 'tumour' of a sort is occasionally encountered. Whitby and Britton/ (1942) refer briefly to rare cases in which chloroma preceded leukaemic manifestations but unfortunately they do not give any details or references. The paradoxical aleukaemic leukaemias are not really additive to the reticulosis hypothesis since so little is known about what determines the entry of the cells into the blood. If leukaemia is regarded as neoplastic then the aleukaemic types or phases might be considered a reasonable parallel to those instances of ordinary malignant disease where metastases at necropsy are either exceedingly restricted or even not demonstrable.

A very rare condition termed histiocytic medullary reticulosis was described by Scott and Robb-Smith, (1939). This was characterised clinically and pathologically by these authors and identified with six similar cases reported previously. This disease appears admirably qualified to be considered a near-perfect example of reticulosis. In the four cases they studied a true generalisation, even to the formation
of macrophage-producing foci in the skin was observed. The perverted hyperactivity of the macrophages (histiocytes) produced an intense anaemia which hastened death in each case. It is possible however to regard the growth as tumour which was lethal by virtue of the erythropagocytosis before it evinced pathological features indicative of that status.

The Papal principle 'quicquid novum, falsum' is usually sound. It must be admitted that reticulosis has been accepted rather uncritically and until recently there has been a tendency to force diseases into the reticulosis schema without adequate justification. As a pathological category it is obviously more elastic than tumour but it is really questionable whether it has deserved its enthusiastic welcome.

Attempts to classify the reticuloses have been many. Four representative schemata by British workers are cited here as they reflect the successive developments of the idea over a period of about 20 years. These works are all readily accessible and need not be repeated, viz. Piney (1925), MacCallum (1928), Ross (1933), and Robb-Smith (1938) and (1944).

It is not proposed to discuss these schemata in detail individually but a few observations are appropriate.

Piney's schema adhered to his concept of the reticulo-endothelial system. The classification was simple and admitted considerable elasticity. The histiocytic (macrophage) scion was not included at all.

MacCallum's classification was on his own admission theoretical. Ross's design was much more an attempt to draw up observed and recorded facts.
facts into a semblance of order and she made a complete distinction between the true and storage reticuloses. It is regrettable that this last feature has not been maintained in later attempts by others. Robb-Smith's system (1938), is widely known, (and also a further elaboration, 1944). One new principle was introduced - the functional medulla. This will be discussed more fully later, but it is equivocal whether it really behaves so independently as the classification suggests.

It has been argued that the more elaborate classifications are too rigid and that like the Ellis classification of Bright's disease they break down at the individual case. This is true but they were primarily an attempt to systematise knowledge out of a chaos of poorly understood entities, and for this they are worthy of much admiration.
TUMOURS OF THE RETICULO-ENDOTHELIAL SYSTEM

This is a difficult and complex subject. The peculiarities of the system in normal growth are retained when it assumes the neoplastic habit. While tumour growth in the tissue may sometimes apparently simplify the problems inherent in it, it more often complicates them in a capricious fashion.

Parts of the system have enjoyed separate anatomical description for a considerable time and even now in systematic studies it is still the practice to treat of these separately. It is convenient to speak of diseases of the blood, diseases of the lymphoid tissue, diseases of the spleen, & etc., but the perpetuation of this method militates against the grander idea of reticulo-endothelial disorders. Notwithstanding, this older method is appropriate in many respects, and within limits emphasizes the weak parts of the system. It has come about therefore to describe tumours of the bone marrow, of lymphoid tissue, of the spleen, and recently of the dermal histiocytes. The representative of the system in the liver is not considered separately, which fact is curious since this part is probably the largest. Exceedingly rare tumours morphologically identical with reticulum cell sarcomata are described from time to time in unorthodox sites e.g. the thyroid gland, testis & etc.

The multiplicity of names given to the tumours of the system is remarkable. This serves to confuse the subject even more. The time-honoured association of botany with anatomy seems to have transferred itself to morbid anatomy, in particular the love of artificial genera. The application of the Natural Orders in classification is only now commencing.
The identity of these growths has often been obscured by well-meant efforts to systematise knowledge by the introduction of better names; which disadvantage is not always off-set by the pictorial presentations of their advocates.

Up till fairly recently tumours of the bone marrow and the lymphoid tissue were recognised in a restricted sense. Newer knowledge with its resultant appreciation of the basis of histogenesis in these tissues has introduced a curious inconsistency which is apparent in retrospect. Thus, while myeloma, chloroma and polycythaemia vera were recognised, entities like diffuse endothelioma of bone (Swing's tumour) and monocytic leukaemia did not enjoy inclusion. In lymphoid tissue the lymphosarcomata have long been known but the tumours more closely related to the reticulum are recent additions. It is singularly unfortunate that this has occurred because it has led to the artificial separation of these newly described neoplasms. In fact, they alone are considered as the tumours of the reticulo-endothelial system in several works. There is not any generally accepted classification yet, but many tentative proposals have been made. Before discussing these, several 'scrap-heaps' require attention.

Most of us are reluctant to abandon the valuable labels round-cell sarcoma and mixed-cell sarcoma. The title, 'round-cell sarcoma', sometimes qualified as large or small, has much to justify its use in actual practice. However, it has now been shewn that true round-cell tumours
rarely arise from mesoblastic tissues other than reticulo-endothelial and visceral muscle derivatives. (Certain authorities e.g. Boyd (1944) include glial tumours). The foremost supply the majority. Accordingly most of these growths are lymphosarcomata and myelomata. In much the same fashion the omnibus 'spindle-cell' and 'mixed-cell' sarcomata have acted as cloaks for tumours, some of which would now be termed reticulum-cell sarcomata. Actually the tumour lymphosarcoma was recognised a considerable time ago, and as far back as 1893 Kundrat sought to differentiate it from Hodgkin's disease; on the other hand the vague round-cell sarcoma was apparently seldom identified with it.

That pathological bete-noir - the endothelial problem - makes its inevitable intrusion. Curiously enough this time the black sheep served to augment the family fortune not a little! In a very careful study in 1913, Ewing produced convincing evidence of the existence of 'endotheliomata' of lymph nodes. In his initial review he referred to several antecedent accounts of 'primary carcinoma' or 'carcinoma alveolare epitheliodes' of these structures. Since reticulum cell growths do sometimes appear epithelial and in contact with one another, these erroneous designations were perfectly understandable mistakes. Ewing was fully aware of the danger of misinterpreting metastatic carcinoma as a primary tumour in a gland and went to great lengths to substantiate that the neoplasms in question were in fact primary. This work was slow to gain credence. It is of interest that Ewing also suggested that the tumours arose from the node reticulum, indeed he even referred to them as reticulum cell sarcomata. This was nine years before Downey (1922)
finally settled the identity of lymph node sinus endothelium with reticulum cells. The concept of reticulum cell sarcoma became further realised. In 1925 Goormaghtigh, in a brief and lucid communication imparted the idea of 'reticulo-endotheliome'. This antedates the widely quoted work of Roulet by five years. His article is illustrated partly in colour which has doubtless aided its popularity, he suggested the term Retothel-sarcom. After this time reticulo-sarcomata were diagnosed so freely that Delbet was prompted to suggest that there was an intoxication in the air responsible!

The status of these tumours is debatable, but when full allowances are made for the extraordinary properties of the component cells of this system, their fantastic interchangeability, their peculiar location and their essential unity, the problem becomes much more straight-forward. Localised tumour is not a necessary qualification for the application of Muir's definition of neoplasm which is adopted. Therefore the leukaemias and polycythaemia vera are properly tumours, the circulatory blood changes merely an inconstant, even incidental, expression of the primary growth in the progenitor cells. Autochthonous growth, the multicentric origin feature which most pathologists emphasize, is nothing more than what is to be expected in a tumour field which happens by nature to be dispersed and worked on a sort of feudal strip system.

The majority of the tumours are malignant. This is peculiar, but not without precedent. Malignant is a relative term and this elasticity is in accord with these growths which show a wide range of behaviour in this respect. This preponderance of malignancy may be related to the essential primitiveness of the cells of the system whereby
even slight augmentation of dedifferentiation at once constitutes frank cytoma. Benign tumours undoubtedly exist; the rare pure myxoma is the best known, even if poorly accredited, example. Stout (1949) is to be congratulated for emphasizing this growth as the important benign tumour of primitive mesenchyme.

Before considering the classifications in more detail it is pertinent to draw attention to the need for certain arbitrary limits to what constitute reticulo-endothelial system tumours. Almost any cell can be phagocytic under appropriate circumstances, but this does not imply that all cells are reticulo-endothelial cells. In the same way connective tissue tumours, e.g. fibroma, chondroma, osteoma, lipoma and chordoma and their malignant counterparts, are not ordinarily included as reticulo-endothelial system tumours. Notwithstanding, the claim of certain fibro-sarcomata and chordomata to inclusion is reasonable, indeed, far more reasonable than the inclusion of histiocytoma (dermatofibroma) which is probably not a true tumour but an hamartoma. If these restrictions are not made then the group becomes unmanageably large.

In this present work emphasis is centred on Hodgkin's disease and its relation to lymphoid tissue tumour. Accordingly these interests receive most attention.

The classification of tumours of the reticulo-endothelial system.

Most schemata suggested present essentially a summary of the growths encountered most frequently. Some workers attempt to qualify these recordings with the clinical aspects, others prefer a purely cellular morphology basis. A few are highly speculative systems of considerable
theoretical interest but like Adami's ingenious lepidoma and holoma theory, extremely difficult to apply in practice. The subject has manifestly proved attractive to numerous pathologists, and it is possible therefore to discuss only a limited number. The examples chosen represent an eclectic series recovered from the writings of the last twenty years. It has already been emphasized that old established portions of the reticulo-endothelial system are often described separately, and certain authors cited below restrict their observations to certain parts. In some cases abstracts from comprehensive writings have been collated for my purpose and this does not imply that the authority from whom the several accounts are drawn supports this.

Ewing (1929) (1941), in his classical text, gives highly critical and profound information on the tumours of the system, but unfortunately it is diffuse and obscure. His classifications are complicated, particularly by the inclusion of granulomata of neoplastic type and the identification of clinical varieties. The full value of his writings is only appreciated after some experience has been gained; they are then recognised as singularly penetrating. It is well known that Ewing retained the term endothelioma despite the difficulties and objections inherent in it. This has caused him to be criticized adversely but it is frequently overlooked that he was far more cognisant of those difficulties than most oncologists. Criticism of the name endothelioma simply because it is no longer fashionable is as unreasonable as criticism of the large lymphocyte of Maximow. Ewing never abandoned the view that Hodgkin's disease was inflammatory and his famous aphorism
'tuberculosis follows Hodgkin's disease like a shadow' reveals where his bias lay. Figuratively Hodgkin's disease can be examined under the enlightening rays of a shadowless lamp, and yet, who can deny that a penumbra lurks is some of the crannies? This concept of neoplastic granuloma pervades his work and has caused great difficulty and misunderstanding. After much thought I have come to interpret it as Ewing's way of stating the very real dilemmas that puzzle honest pathologists, especially those who have studied many doubtful lymph nodes, namely, is it inflammatory?, is it tumour?

The classifications he offered were tentative. In lymphoid tissue tumour he considered that the lymphocyte might give rise to a lymphocytopma which may take the form of a benign circumscribed growth, in certain cases initiated by tuberculous infection, a malignant lymphocytopma or lymphatic leukaemia, and pseudo-leukaemia (probably Hodgkin's disease is meant). The reticulum cells of the same tissue were credited with giving rise to large round-cell hyperplasia or neoplasia expressed as a granuloma malignum, myeloid leukaemia, large cell lymphosarcoma or Hodgkin's sarcoma. The endothelial cells were accorded specificity in this schema and their overgrowth or neoplasia acknowledged separately. It should be noted, however, that in the relevant text and in his 1913 paper, the identity of lymph node reticulum cells (Ribbert's type) and lymphatic endothelium are virtually admitted. His rare diffuse endothelioma of bone (endothelial myeloma) corresponds to reticulum cell sarcoma in that site.

Warthin (1931) did not actually submit a classification (he had /already
already done so in 1904), but this work is highly relevant as it is the first reasoned discussion on the generic affinities of Hodgkin's disease and mutations between certain lymphoid tissue tumours. The hypothesis that Hodgkin's disease is neoplastic was well propounded, and a full descriptive account of the morphology of Hodgkin's sarcoma was given. The triad, Hodgkin's disease, lymphosarcoma and the less well characterized mycosis fungoides, were shewn to merge imperceptibly into one another. These transitions are to be sought in the study of examples of these diseases which present in an atypical form. This communication is deficient in actual presentation of illustrated instances, but it is offered by an undoubted authority.

Callender (1934) produced a comprehensive integrated system of classification of the tumours of the reticulo-endothelial system. This was based on a complex analysis of 380 cases collected over a period of seven years by the Lymphatic Tumour Registry. The classification is derived from the presumed progenitor cells of lymphocytes, granular leucocytes, erythrocytes and monocytes. The first category contains lymphatic leukaemia, lymphatic pseudo-leukaemia and lymphosarcoma. Callender observed that true lymphoma, meaning a benign circumscribed tumour of lymphocytes, was not represented and probably did not exist; accordingly he held that it was illogical to call the malignant counterpart lymphoblastoma. For this reason he equates lymphatic leukaemia with 'lymphocytoma, leukaemic'. His lymphatic pseudoleukaemia is equivalent to 'lymphocytoma, aleukaemic'. This corresponds in his view to two diseases viz. aleukaemic lymphatic leukaemia (diffuse lymphatic
pseudoleukaemia) and giant follicular hyperplasia with splenomegaly, Brill's disease, (nodular lymphatic pseudoleukaemia). Lymphosarcoma in this schema is characterized by the development of true metastases and is typically aleukaemic. The latter feature may be lost in some cases.

The characterisation of the varieties listed is very good and full appreciation of their lability is acknowledged. On the other hand, criticism of the nomenclature adopted is justified, particularly the use of pseudoleukaemia which has long been a synonym for Hodgkin's disease. In addition, the clinical features which are itemised for the various forms are insufficiently stable to reinforce the subdivisions very much.

Under the granular leucocyte heading are listed leucocytosis, myeloid leukaemia, aleukaemic myelocytoma and myelosarcoma. Myeloid leukaemia is termed 'myelocytoma, leukaemic', the aleukaemic form being identified with single or multiple myeloma. Since Callender agrees that the commonest form of the latter is the plasma cell type, the propriety of including this tumour here is open to question. Myelosarcoma is recorded as possibly occurring in an aleukaemic form and in a leukaemic form which is also termed myelocytic leukosarcoma. These categories include the green tumour, chloroma. Callender believes that the members of this group are more stable entities than the lymphocyte tumours and that the most difficult problem is the establishment of satisfactory criteria to recognise sarcomatous change among them. This however, he admits, is not of much practical importance.

The erythrocyte series is represented by the polycythaemias including the symptomatic forms. Polycythaemia vera is considered /dubiously
dubiously homologous to leukaemia and the doubtful erythrocytoma is mentioned.

The last group is collected under the heading 'Reticulocyte'. The usage is confusing, the reticulum cell or monocyte being better equivalents. The monocytes of specific infections and the storage diseases are classed together as reactive reticulum cell hyperplasias. Under the sub-heading of reticulocytoma, leukaemic and aleukaemic varieties are described. The first corresponds to monocytic leukaemia. It is to be noted that Callender manifestly leans to Naegli's interpretation of this disease, the Schilling type is only doubtfully acknowledged. The aleukaemic form is equivalent to the reticulum cell sarcoma of other authors. This latter identification is amongst the first statements of this idea.

Hodgkin's disease is considered as belonging to a special category but still under the general reticulocyte heading. Two forms are described, one corresponding to the classical malady and another apparently the counterpart to acute Hodgkin's disease. Hodgkin's sarcoma is a further special category, being regarded as a reticulum cell sarcoma which is characterized by an high degree of pleiomorphism. In his notes on this section, Callender criticises the view that transition occurs as much as is suggested by some writers, e.g. Herbut, Warthin & etc., and makes a plea for careful studies of the natural evolution of diseases in the group. The schema has been criticised widely but it is a very considered and thoughtful exposition.

Ehrlich and Gerber (1935), studied a series of 18 cases of lympho-
sarcomatosis in considerable detail. The examples were examined fully at necropsy and most of these patients had had antecedent biopsies performed. These workers decided that the tumours fell into three main categories using as a basis for this separation the morphological characteristics of the predominating cell type. These cell types were found to correspond in their essential morphology to the immature, intermediate and mature cells resulting from abnormal differentiation of the cytoplasmic reticulum along lymphopoietic lines. This hypothesis of reticulum cells giving rise to lymphatic series cells is one of the main contentions of these authors, and they frequently invoke the authoritative work of Klemperer in support of it. So far as I am aware, this is the first acknowledgment of the 'bipotential stem-cell' idea in reticulo-endothelial system tumour. They classified their tumour as follows:—

1. Cases in which large pale cells in reticular arrangement predominated. Formation of new reticulin fibrils was minimal or absent. This variety they termed 'reticular'. It appears to correspond with 'stem-cell lymphoma' and 'lymphoblastic lymphosarcoma'.

2. Intermediate type. A group in which mixed cells occur, partly reticular (syncytial) and partly free. Some new reticulin formation was characteristic.

3. Lymphocytic type. A group in which the tumours were composed predominantly of free cells either of the immature or mature lymphocytic type. This growth seems to correspond to the lympho-
cytic reticulum cell sarcoma of Robb-Smith (1938) but they add that new reticulin is formed in considerable amount. It is this last feature which is difficult to accede to readily if the neutral path of orthodoxy is followed.

This subdivision does reflect what occurs – the need for the recognition of intermediate mixed reticulum cell/lymphocytic cell growth is very real.

In their discussion the mode of growth is fully and, I believe, very ably presented, in particular their views regarding partial – almost focal – involvement of lymph nodes in early stages are most welcome. One interesting peculiarity is the rarity with which their cases exhibited splenic involvement (2 cases); this they observe may be related to the fact that this organ is not true lymphoid tissue since it lacks lymphatics. Limited systematisation of the process is another feature they emphasize and contend that this is in contrast to hyperplasia ('lymphadenosis'). Their views are extended to Hodgkin's disease in which they hazard that both fibroblastic and lymphopoietic potentialities are developed, the former being the major expression. Brill's disease they regarded as a link between lymphadenosis and lymphosarcomatosis.

Dawson, Innes & Harvey (1937), in the second of their admirable "Debatable Tumours" series discuss the same problem in detail. They define their subject as, 'lymphosarcoma or reticulum cell sarcoma, a radio-sensitive tumour of lymphoid tissue arising from the embryonal stem cells of that tissue'. They state that in the fully developed tumour the normal micro-architecture of the lymphoid tissue is lost, and
the cellular appearance becomes more uniform. The tumour cells are invariably larger than small lymphocytes according to these authors, (correspondingly they reject 'lymphocytic reticulum cell sarcoma' as a true neoplasm). In form the cells may be polyhedral, ovoid or spheroidal with very variable cytoplasm content. Mitoses are always to be found. The nucleus is described as somewhat vesicular and containing more chromatin than the parent reticulum cells. Characteristically the tumours are locally invasive and though some deposits are undoubtedly autochthonous, blood borne metastases are probable.

The histogenesis is discussed very fully on the basis of their summary of normal lymph node structure which is a particularly lucid and acceptable account. They give the following schema indicating the relationships between the cells of the reticulo-endothelial system:

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Primitive mesenchyme cells.
  fibroblast  reticuloblast  lymphoblast
  fibrocyte   macrophage   littoral endothelial lymphocyte cell
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and suggest that 'mesenchymal' should be substituted for 'reticulo-endothelial'. This might have advantages, but the former word is used in different senses and the idea is too late. The bi-potential stem-cell idea is fully endorsed; they write 'reticulum cell and lymphopoietic lymphoblast we consider the same cell or at least only slightly separated from the stem cell'. The illustrations though small, are excellent.

Towards the end of the paper Hodgkin's disease is discussed and /curiously
curiously treated. It is insisted that a sharp distinction is to be made between the two lesions yet at the same time it is admitted that this is constantly a source of difficulty. Whereas in the earlier part of the study, multinucleated giant cells, which answer perfectly to those of Hodgkin's disease, are described as occasionally to be seen in their reticulum cell sarcoma; and fibrosis, represented by variable reticulin increase is held to be common, they seem to recant suddenly and by-pass the only logical solution: Namely, they are variants of the same process. In fairness it is only right to the authors to recognise that they did not set out to settle this last question. This is an highly instructive communication and establishes the generic reticulum cell sarcoma beyond reasonable doubt.

Robb-Smith (1938) put the whole reticulosis and reticulo-sarcoma problem on a sophisticated basis which has rightly prompted universal admiration. Reticulosis has already been discussed and here it will suffice to consider the latter. At the outset of the second part, this author stresses the peculiar difficulties attendant on distinguishing true tumour from primary reticulosis and advances the phenomenon of stromal destruction as the only reliable criterion. In the classification of the true reticulo-sarcomata in lymph nodes he all but abandons the follicular, sinus and medullary theme and reverts to a cytological basis. The schema resulting introduced a much more comprehensive concept of reticulo-sarcoma than had been contemplated previously, or at least integrated by earlier workers.

/Reticulo-sarcoma.
Reticulo-sarcoma.

A. Undifferentiated reticulo-sarcoma (syncytial).
   1. Diffuse.
   2. Trabecular (stroma reaction).

B. Differentiation to histioid cells.
   1. Dictyosyncytial (fibrillo-syncytial) reticulo-sarcoma.
   2. Dictyocytic (fibrillary) reticulo-sarcoma.

C. Differentiation to haemic cells.
   1. Lymphocytoma.
      (i) Lymphoblastic sarcoma.
         (a) Medullary.
         (b) Follicular.
      (ii) Lymphosarcoma.
   2. Plasmocytoma.
   3. (Monocytoma).
   4. Myeloblastoma.
   5. Erythroblastoma.

D. Mixed type (polymorphic reticulo-sarcoma).

E. Differentiation of sinus lining cells.
   (a) Undifferentiated cell type (reticulo-endothelio-sarcoma).
   (b) Differentiated cell type (histiocytoma).

The growths classified in A. and B. are essentially variants which are very closely related, the distinction probably not being of very deep significance except theoretically.

C.1(i) corresponds to the familiar large cell lymphosarcoma, the sub-divisions
sub-divisions (a) and (b) connote the type of origin or presumed origin; (b) indicating development from a previous Brill's disease. C.1(ii) is the small cell lymphosarcoma - the growth to which Dawson et al (1937) take exception. The remainder, C.2., C.3., C.4., and C.5., since the classification is primarily descriptive of lymph node tumours, are rather hypothetical. The author does not discuss them beyond a brief note. The omission of C.3 is interesting since Callender (1934) considered monocytic leukaemia as a reticulum cell sarcoma of lymph nodes, etc.

Category D., Robb-Smith holds, corresponds to many cases of Hodgkin's sarcoma or malignant Hodgkin's disease. This question will receive further attention later.

Category E. is again chiefly included for completeness. (The omission of myxoma is curious considering that histiocytoma is included).

This schema is open to many criticisms when tried out in practice. A common observation is that several variants may often be found in one and the same case, or even node. On the other hand the classification is only what it is stated to be - a histological one, it does not pretend to be a cut and dried résumé of what actually happens. At the same time it is not entirely speculative and is of extreme theoretical interest because it has a catholicity fully reflecting the potentialities of reticulo-endothelial system cells.

In 1941 Warren and Picena published a valuable contribution to the subject of reticulum cell sarcoma in lymph nodes. In particular they gave an excellent critical review of the criteria advanced from time to
time to establish the identity of these tumours and gave it as their opinion that the diagnosis of reticulum-cell sarcoma was probably too readily and too frequently made. The writers adhere to Oberling's definition of reticulo-sarcoma indifferencié (1922). This definition is considered rather restricted by some workers but is very helpful nevertheless. 'The new growth is formed of syncytial masses of an undivided or slightly fenestrated protoplasm whose limits are ill-defined and connected with zones of more or less differentiated lymphoid tissue. The fenestration, by exaggerating the internuclear spaces, may give a reticular structure. In these undivided masses of protoplasm there are many irregularly distributed oval or indented nuclei, with granules, and one or two prominent nucleoli. The nuclei often appear almost empty. Mitoses are observed and some isolated tumour giant cells are common. It is not rare to see some lymphocytes scattered through the syncytium'.

In their own studies these workers record very variable impregnation results and significantly they set little store on its value. They suggest the following classification:

1. **Reticulum cell type** - undifferentiated or syncytial.
   - reticular (dictyocytic).
2. **Mixed type** - combination of reticulum cells and lymphoid cells.
3. **Lymphoblastic** - large cell (lymphoblastic lymphosarcoma).
4. **Lymphocytic** - small cell (lymphocytic lymphosarcoma).

In so far as it goes this classification is good because these /variants
variants do really happen, in particular their category 'mixed type' is welcome. In their discussion they state 'our concept of the normal lymphoid tissue is a network of undifferentiated cellular reticulum (stem tissue) in whose interstices lie lymphoid cells; (lymphoblasts, prolymphocytes and lymphocytes), the offspring of the former. Thus, the complex of reticulum cells and lymphoid cells cannot be destroyed without destroying the concept of the essential unity of lymphoid tissue. Proliferation of one almost invariably causes variations in the other'.

They give several very convincing, well illustrated examples bearing out the truth of this belief and include an ingenious drawing depicting the subtle transition of predominant reticulum cell neoplasm to predominant small cell lymphosarcoma.

Although restricted to lymph node growth, the essential unity of lymphoid tissue is emphasized and the generic lymphoid tissue sarcoma is clearly envisaged in this paper.

Gall and Mallory, (1942), give a discursive and exhaustive analytical account of a series of lymphoid tissue sarcomata (malignant lymphoma) extending to 618 cases. They uphold the cytological basis of classification exclusively and criticise Callender adversely. Their definition of malignant lymphoma, 'a generic term to designate those maladies which are characterised clinically by progressive tumour-like enlargement of lymphoid tissue with eventual death', is interesting as it covers the reticuloses. This latter concept does not appear to have gained much approbation in the U.S.A.

In their studies they concluded that these growths could almost all
be classified as follows:-

1. Where the histological pattern is simple. (This appears almost equivalent to one-cell-type tumours).

(a) Stem cell lymphoma. 
   Incidence. 56 cases

(b) Clasmatocytic lymphoma.
   " 71 "

(c) Lymphoblastic lymphoma.
   " 85 "

(d) Lymphocytic lymphoma
   " 135 "

2. Where the histological pattern is complex.

(e) Hodgkin's lymphoma.
   " 193 "

(f) Hodgkin's sarcoma.
   " 36 "

(g) Follicular lymphoma.
   " 42 "

Total 618

Note: Stem cell lymphoma is the undifferentiated reticulum cell sarcoma of other writers; the clasmatocytic growth is a tumour of cells morphologically equivalent to monocytes and macrophages. In subdivision 2, Hodgkin's lymphoma is ordinary Hodgkin's disease.

It is notable that all of the series could be fitted into one of the seven different categories. This postulates a definite elasticity or their series must be unique; I collected only 220 and in finer details anyway, I could devise almost as many categories!

The most significant aspect of this classification is the inclusion of Hodgkin's disease. As will be seen later, the view that Hodgkin's disease is tumour is as old as its discoverer and therefore no new thing, but here it is openly included in a schema of lymphoid tissue sarcoma for the first time. Another important view of these authors is that within the categories above there is great constancy of morphology and whilst admitting variations do occur, they insist that these are practically /limited
limited to degree of differentiation without significant transition between one major group and the other. In 84 of their cases either repeated biopsies or subsequent necropsy, or both, were carried out. In 56 of these patients there was an interval of at least one month between the procedures. Analysis yielded the following results:—

Please see over-leaf.
<table>
<thead>
<tr>
<th>Condition</th>
<th>Average interval between specimens (years)</th>
<th>Cases</th>
<th>Unchanged.</th>
<th>Dedifferentiation.</th>
<th>Differentiation.</th>
</tr>
</thead>
<tbody>
<tr>
<td>Stem cell lymphoma.</td>
<td>0.25</td>
<td>2</td>
<td>2</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Clasmatocytic lymphoma.</td>
<td>2.30</td>
<td>12</td>
<td>6</td>
<td>6</td>
<td>-</td>
</tr>
<tr>
<td>Lymphoblastic lymphoma.</td>
<td>0.75</td>
<td>4</td>
<td>2</td>
<td>2</td>
<td>-</td>
</tr>
<tr>
<td>Lymphocytic lymphoma.</td>
<td>1.90</td>
<td>9</td>
<td>9</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Hodgkin's lymphoma.</td>
<td>1.90</td>
<td>20</td>
<td>16</td>
<td>4</td>
<td>-</td>
</tr>
<tr>
<td>Hodgkin's sarcoma.</td>
<td>0.5</td>
<td>1</td>
<td>1</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Follicular lymphoma.</td>
<td>5.0</td>
<td>8</td>
<td>7</td>
<td>1</td>
<td>-</td>
</tr>
<tr>
<td><strong>Total</strong></td>
<td></td>
<td><strong>56</strong></td>
<td><strong>43</strong></td>
<td><strong>13</strong></td>
<td>-</td>
</tr>
</tbody>
</table>
It is seen that 43 (76.82 per cent) retained the original structure and 13 (23.2 per cent) shewed dedifferentiation. It is noteworthy that all the alterations recorded were atavistic. This latter figure is too large to be dismissed lightly and lends support to the view that the growths are potentially labile phases of lymphoid tissue sarcoma and not fixed entities.

Except for refinements or broad generalisations, the classification of lymphoid tissue tumours has not been altered much in the last decade. The unity has been fairly generally accepted, Hodgkin's disease and Brill's disease have been included, and a degree of transition from one morphological variant to another has been admitted. It is concerning the last phenomenon that appreciable advance has been made in this period.

Custer and Bernhard, (1948), reinforced this last belief. Their paper includes an excellent critical account of the gradual development of the hypothesis that Hodgkin's disease is neoplastic and a comprehensive study of the mutations encountered in a very large series of cases of lymph node tumour. Their series consisted of 200 necropsy cases, 138 of which had had antecedent biopsies; together with 500 biopsy examples. In the latter material 431 were single specimens, 60 were two specimens at intervals, and 9 were three specimens at intervals. 33 per cent of the total cases studied shewed alterations in the course of their natural history. The following schema indicates the nature of the alterations observed:-

/Follicular
Their experience determined that follicular lymphoblastoma was the most labile variant of the group. They were unable to confirm that X-ray treatment had any influence in causing the transformations noted. The illustrations to this paper are copious and excellent.

It is overwhelmingly convincing that so far as the variants scheduled are concerned, there is not any doubt whatever that lymphoid tissue tumour is to be regarded as a single neoplastic entity which may be expressed in different histological patterns. The one regrettable feature of this work is the absence of full characterisation for the diagnosis of Hodgkin's paragranuloma and Hodgkin's sarcoma.

Willis (1948) in a masterly chapter simplifies the problem of lymphoid tissue tumour in an emphatically practical manner. In essence his views include a central concept of lymphoid tissue sarcoma, which tumour conforms broadly to one of four main morphological expressions. His schema is as follows:

```
Follicular Lymphoblastoma


Mutation in indicated direction only.

Mutation in either direction, occasional.

Mutation in either direction, frequent.
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/Lymphoid
Lymphoid tissue sarcoma.

Main variants:

(a) Follicular lymphoma.
(b) Lymphosarcoma, with or without leukaemia.
(c) Hodgkin's disease.
(d) Reticulum cell sarcoma.

A simple classification admits of much elasticity and it is partly for this very reason that Willis prefers it. The previous more complex schemata tended to emphasize unduly the expressions of lymphoid tissue tumour as entities, which is misleading. On the other hand they exemplify the most praiseworthy endeavours which must of necessity precede the broad generalisations - namely the study of things caused.
PART II

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THE LYMPHATIC SYSTEM

Foreword:

This body system has long enjoyed the distinction of separate consideration in Anatomy. Despite the advent of its parent, the reticulo-endothelial system, it is still common practice to accord it separate attention. The selection of a portion of a diffuse and complex system suggests that simplification would result but this unfortunately does not follow. The reasons include inseparability of sibling from sire and a failure to define the lymphatic system satisfactorily.

In most standard anatomical texts the definitions of the lymphatic system are brief and insufficiently comprehensive. Le Gros Clark (1945) acknowledging the deficiencies, defines it thus:— 'The lymphatic system in its narrower sense consists of a system of fine absorptive vessels permeating most of the tissues of the body and emptying their contents ultimately into the venous system. Along the course of these vessels are groups of lymph nodes ...... localised masses of lymphoid tissue of different types closely related in structure to these are also to be found in other parts of the body ....'

It is proposed to offer some account of the system as defined by Le Gros Clark and to augment this by observations on the human lymph nodes.
THE LYMPHATICS
with observations on some of their functions.

The structure of the lymphatic vessels is simple. The typical lymphatic capillary has a wall made up of a single layer of endothelial plates. The lymphatic trunks are lined by a similar intima surrounded by scanty connective tissue. Smooth muscle is interposed between these tunicae in the largest trunks but rarely forms a perfect media. In all animals which do not possess lymph hearts the lymphatic trunks and larger lymphatics are valved, and these structures follow one another at much more frequent intervals than in veins.

Lymphatic vessels are not present in tissues which are devoid of a blood supply; namely epidermis, hyaline cartilage, nail, enamel, and the cornea. One further certain exception is the central nervous system, despite its high vascularity. It has been suggested by Woolard (1924), that the Virchow–Robin spaces act as lymphatics in this situation. In support of this view, which incidentally implies the assignment of lymph-like properties to the cerebrospinal fluid, is the fact that it drains into the venous system via the Pacchionian bodies. On the other hand pathological increase of protein in the fluid is apparently reduced very slowly. This phenomenon is well seen in patients recovering from anterior poliomyelitis where high protein levels persist long after the pleocytosis has regressed. In ordinary lymphatics the protein level is hardly ever more than half the blood protein level.

It is now established that the serous cavities, the pleural and peritoneal sacs, joints and bursae are not part of the lymph vascular system as von Recklinghausen believed. Representatives of the
lymphatics are very doubtfully present in bone and muscle apart from the perivascular channels. In certain internal organs lymphatic vessels have a restricted distribution. In the liver they are confined to the portal canals and in the lungs to the immediate vicinity of the bronchial arborisation. Similarly in the spleen lymphatics are confined to the sheaths of blood vessels in the trabeculae, and then only in the region of the hilar intrusion.

Elsewhere they are disposed in superficial and deep plexuses. The former are exceedingly large and are related particularly to surfaces, where they form vast mutually intercommunicating networks almost comparable in extent to the blood capillary mesh.

The anatomical discovery of this inconspicuous vascular system was accomplished over 300 years ago. The lymphatic vessels were first observed in 1627 by Gasparo Asellius, Professor of Anatomy at the University of Padua, who noted them in the mesentery of the intestine of a well-fed dog. He also appreciated their association with the mesenteric lymph nodes which in this animal as in the cat and rabbit, form a tolerably discrete conglomerate mass, and are appropriately known as Asellius' pancreas. (The purist may complain that a second pancreas is a bad idea to perpetuate, but after all the word means 'all meat', and is not so very apposite for the sweetbread which in man very commonly shews some degree of fat replacement). It is noteworthy that their discovery was related to their being filled with opaque chyle at the time of the historic observation. It seems fit to record in honour of Asellius that this well known and undeniable task performed by the /lymphatics,
lymphatics, viz., the absorption of fat from the intestine is still an almost solitary agreed fact. This property does not appear at first to have much connection with what may be their function elsewhere, nevertheless there is one small point of comparison which is seldom considered. This concerns the dimensions of the fat particles absorbed. Although this subject appears a dangerously physiological problem for one schooled to think in terms of dead issues, courage is gained from the following statement. 'The mechanism of fat absorption is a highly controversial subject' Samson Wright (1945). A frank admission of this order from authority is tantamount to saying that one guess is as good as another. The proposal that fat is directly absorbed in very highly divided form is attracting serious attention once more. The original observation of Krehl in 1890 - when osmic acid ($\text{OsO}_4$) enjoyed a higher reputation for specificity than it does now - purported to shew how exceedingly minute droplets of fat appeared in the intestinal epithelial cell cytoplasm, and by coalescence formed larger globules. These then made their way into the villus lacteal, the transit being effected by leucocytic activity. This latter has long been a disputed mechanism and still merits the good Scottish verdict of Not Proven. This theory fell into disrepute after it was decided that hydrolysis of fat was the initial step in the absorption process. Later when it became popular to visualise the fatty acids as little fir-tree-like molecules in which only the 'root-portion' - the carboxyl group was soluble, the problem of their entrance had still to be considered. The configuration of the long chain fatty acids does not appear conducive to an easy passage, even with lecithin
as a lubricant. Molecular distortion in proteins is an established phenomenon but is not reported to occur in fatty acids. The emulsion of fat in the upper small intestine reduces the particles to chylomicra which have been estimated by Gage and Fish (1924-5) to be 0.5 - 1.0 μ (500 - 1000 μ) diameter. These dimensions are well within the scope of ordinary microscopical method but in terms of physical chemistry such particles are enormous. It is probable that the fat elements that actually traverse the villus cell wall are much smaller. With a good microscope it is possible to see but not resolve, particles as small as 0.074 μ (74 μ). (They can indeed be resolved by Barnard's ultraviolet light microscope, and of course offer little theoretical difficulty to fuller examination by electron microscopy). In dark ground examination of a fine emulsion e.g. milk, it is possible to see particles which are probably fatty which are much smaller than 0.5 - 1.0 μ, and furthermore it is probable they are spherical. If good histological preparations of intestine during fat absorption fixed with osmium tetroxide are examined with the oil immersion objective particles of osmophile material well below 1.0 μ diameter are discernible.

Elkes, Frazer, and Stewart (1938), made measurements of chylomicra in human blood and reported that the majority lay in three ranges of the orders of 1μ, 1/3 μ and 35 μ. They also considered that all the evidence available supported the belief that the particles in question were fat. Frazer (1946) in a review on the theories of fat absorption, maintained that provided emulsification was sufficiently complete even paraffin was absorbed. This fact which is capable of demonstration
indicates that particle size is almost certainly an operative factor.

The evidence for the existence of exceedingly finely divided fat is reasonably convincing and this brings the phenomenon of fat absorption by the intestinal lymphatics nearer to the functions of other lymphatics than is at first obvious. This connection relates to certain aspects of protein metabolism in which the lymphatic system is concerned. The link lies in the reasonable comparability of particle size between the ultimate forms of fats and proteins. The size of protein molecules is of an extremely small order. Lloyd and Shore (1938) have abstracted a large series of papers relating to protein molecular size expressed in absolute terms. The molecules of pepsin, trypsin, ovalbumin, insulin and muscle haemoglobin are reported to be spheroidal with diameters of 4.4 μ. Haemoglobin and serum albumin have not spheroidal molecules but calculated as spheres they would be 6.8 μ. It is further stated that one milligram of protein (type not specified) will present an area of one square metre. This has been calculated to correspond to the surface area presented by a set of spheres of radius about 2.2 μ. Svedberg (1937) found that most proteins form homodispersed solutions with molecules which are spheroidal or at least approach that form.

Regarding their size and molecular weight he decided that they fall into two classes. A group with a diameter of 2-4 μ with molecular weights approximating to a multiple of 34-36,000 and a group of 72 μ diameter with molecular weights of the order 400,000. It is widely agreed that natural proteins usually exist in multimolecular aggregates, which postulate is in accord with their colloid properties. Thus there is
some justification for believing that no great disparity exists between the fat and protein particles at certain metabolic levels.

The lymphatics are concerned with two main functions relating to the transit of protein in the body. The first pertains to the conservation of the integrity of the vascular system and the second to defence mechanisms of the body.

While ideas concerning the functions of the lymphatic stream generally are still relatively meagre, it has gradually come to be postulated that it is a vascular contrivance which off-sets certain defects of blood vascular systems. This role is emphasized by Drinker and Yoffee, (1941) and the concept is reinforced by Le Gros Clark (1949). In essence this function is the transport of extravasated protein back to the blood. In animals with an highly developed blood vascular system the blood stream subserves the majority of the metabolic functions. In the process of nutrition food substances and oxygen are yielded to the tissues at capillary level. The gaseous exchanges are not relevant here, but other metabolites in dialysable form traverse the capillary impelled by osmotic and hydrostatic mechanisms. It appears that minute amounts of plasma protein inevitably leak and the raised colloid osmotic pressure at the venous end of the capillaries is not sufficiently powerful to recapture it. It is believed that this material and fluid - sufficient to retain tissue-fluid at an inconsiderable level - are absorbed by the lymphatics. Thus in one view the system may be regarded as a compensatory apparatus to correct deficiencies which are inherent in blood vascular systems utilizing high pressure. Phylo-
genetically lymphatic systems are old but blood vascular systems are older so this idea is reasonable.

In general terms there is evidence that the working pressure of the blood vascular system rises as the phylogenetic scale is ascended, and concurrently the lymphatic system becomes more complex and efficient. The carriage of protein by the lymph is well established fact yielded by direct examination of the fluid in efferent trunks in experimental animals. This is readily performed in the frog which has easily accessible lymph hearts. In birds and mammals it is a protracted and difficult procedure calling for the highest skill and patience. It entails not only identifying the lymph trunk chosen but cannulating it, initiating a lymph flow in it, and maintaining that flow long enough to serve the purpose.

In man comparable experiments are not applicable and opportunities to study similar circumstances arising from injury or disease are rare. Drinker (1946) in a most thoughtful and delightful contribution to a symposium on lymph, draws attention to the following remarkable instance which is fully recorded by Crandall, Barker and Graham (1943)*

A negress aged 30 years sustained a gun-shot wound of her neck whereby her thoracic duct was severed. The lymph fistula resulting did not heal spontaneously, an unusual event, and a careful study was made of its discharge (lymph), and the blood, together with recordings of the patient's weight over a period of six weeks. It was found that a weight loss of 5 lb. a week resulted and that the plasma proteins fell to 3.5 gm. per cent in a month. High protein feeding resulted in a slight
rise of plasma protein (to 4.6 gm. per cent in thirteen days). At this point the thoracic duct was ligated and the patient began to regain weight. Drinker points out how with the lymph lost to the blood a progressive depletion of protein resulted which unchecked would probably have been fatal.

I have personal experience of a similar case. During the assault on Sicily in 1943 the Unit I served in received casualties by air from the beaches. A young German paratrooper who had been repeatedly bayonetted in the chest was admitted within 12 hours of wounding. He came under the care of Lt/Col. d'Abreu R.A.M.C. who established the diagnosis of haemothorax. Paracentesis thoracis revealed several pints of blood in the left chest which were drained and the patient was treated by blood transfusion. The following day signs of fluid had reappeared in the left chest and paracentesis disclosed approximately seven pints of pale pink milky fluid. This drainage alleviated his discomfort very considerably. The fluid was shewn to contain fat and formed a gel on standing, protein estimations however were not done. It remained sterile on culture. It was found necessary to aspirate the chest daily to combat cyanosis and embarrassment. His general condition deteriorated and he became oedematous and died about ten days after admission. At necropsy healing wounds of the lung and liver were confirmed and the thoracic duct was found to be severed; the cut ends were retracted and everted recalling the appearances of bowel wounds when missiles traverse them. The injury had been caused by a glancing blow of the weapon striking the spine. In this case it is doubtful whether
lymph loss alone accounted for death since the duration was brief, clinically the mechanical effects of the fluid appeared the chief derangement operative.

At this point it is appropriate to allude to the extraordinary mystery of how the lymphatics work, and what they are capable of accomplishing.

In the frog the lymphatic system is a conspicuous fraction of the vascular apparatus and it has been studied extensively. In this animal the system is not closed but communicates by means of stomata with the tissue spaces. An anecdotal and amusing fact with regard to these stomata is recorded by Klein. The cells lining them become ciliated in the winter time - but only in the female of the species! These animals have in addition lymph hearts, four in number. Thus, in the female frog cilia at the radicles of the system, and hearts in the course of it, serve to propel the lymph back to the blood, in the winter; presumably lacking the ciliary aid in the summer. This naïve account is not offered very seriously but a source of motive force is at least apparent in it, and inductive portals of entry are exhibited.

In the higher vertebrates, including man, it is very generally agreed that the developed lymphatic system is a closed one, Drinker and Yoffe (1941) and Le Gros Clark (1945). Further, lymph hearts do not exist. The lacteals are the only lymphatics in man which show typical contractility. Yet material of relatively enormous molecular weight effects entry readily via non-alimentary lymphatics. Barns and Trueta/(1941),
69.

(1941), investigating the route of absorption of venom and toxins found that Black Tiger snake venom, molecular weight over 20,000 and Russel viper venom, molecular weight at least 30,000, were almost entirely absorbed by lymphatics. Diphtheria toxin with a molecular weight about 70,000 was also absorbed exclusively by these channels. This latter is an important observation since plasma albumin has practically the same molecular weight (67,500 - 70,000+), Harrison (1934). Cobra venom with a molecular weight between 2500 and 4000 enters the blood stream directly at about the same rate as via the lymphatics.

Exactly how this 'absorption' is effected is obscure. The common term applied to the lymphatic channels of a part, namely the lymph drainage, suggests a sophisticated sewage system with the thoracic duct naught else but a thinly disguised Cloaca Maxima. Admittedly the pressure at this Ostia is low, probably in the order of 5-15 cm. water, but it is not much lower than the blood capillary pressure, Wright (1945).

In a patient with general chronic venous congestion this pressure (capillary) may rise to twice the latter figure, i.e. 30 cms. water, but lymph still drains. If it did not he would assuredly die, as a frog does when its lymphatics are obstructed, and it is a fact that in cases of the above nature protracted chronic dropsy is not uncommon. This diminishes the strength of the argument which assumes that a pressure gradient is responsible for the shift. In any event it is decidedly peculiar for the effluent of drains to proceed uphill! The presence of valves in the lymph channels (beyond the capillaries) stimulates the imagination and one might envisage a kind of hydraulic ram mechanism /effecting
effecting the lift of lymph from say the foot to the shoulder. This however is a proposition with little to support it — where is such an intermittent force to come from? The arterial pulse does not extend down to the true capillaries and the rhythmical opening and closing of these channels is a gentle affair.

In experimental studies of lymph production in mammals it is invariably difficult to obtain a flow in the cannulated lymph trunks of the anaesthetized animal. The flow can be augmented by producing passive movements in the beast and ingenious mechanical devices to effect this have been elaborated. As a corollary it is supposed that muscle movement is a potent factor in promoting lymph flow.

This factor is undoubtedly important particularly in visceral muscle activity. It seems conceivable that the lethal effects of paralytic ileus are related to the lymph stasis resulting. On the other hand patients with paraplegia or bed-ridden people do not become oedematous by reason of muscle inactivity alone. The invocation of incidental muscle movement as a factor in the transport of body fluids is ingenious but suspect. In the case of venous return it is merely subsidiary, the left ventricle effecting most of the shift.

The above remarks do not much contribute real understanding of the crucial problem, namely how do such enormous particles get into the closed lymphatic. It has been shewn, Clark and Clark (1926), that a poor kind of phagocytosis is a property of lymphatic endothelium. This however can be rejected here because the endothelium studied was recently repaired and the process was very slow. Personally I find the /situation
situation quite incomprehensible. The entry of microbes does not present so much difficulty if the generally accredited mechanism is the only one. It assumes that the micro-organisms are ingested first by microphages or macrophages and that these cells then enter the lymphatic in the same mysterious fashion in which they enter or leave blood capillaries. The entry of viruses is in all probability direct. In most cases their dimensions, 10 - 300 μ, are conducive to belief in this hypothesis.

Note on the Development of the Lymphatic System

This subject has received little attention in the study of human embryology, most of the restricted work performed has been conducted on the lower animals, notably the chick and the pig.

Frazer, (1931), states that lymphatics first appear in the human embryo at or just after the 10 mm. stage, i.e. in the sixth week of intra-uterine life. The origin of the lymphatic vessels is still a disputed problem. Two main views have been held, both ably supported by competent observers, and both open to certain criticisms. The one idea is that the lymphatics arise from veins, and the other that they arise de novo in the mesenchyme. The former view is more favourably regarded, Le Gros Clark, (1945). In this connection it is significant that permanent intercommunication between the lymphatic vessels and the veins is constantly established at the terminations of the thoracic and right lymphatic ducts. It is difficult to imagine why an highly complex subsidiary vascular system of lymphatics should come into being independently and then acquire anastomoses at these curious loci with
such regularity. On the other hand the arguments of Huntingdon (1914), and critical support of McClure, (1951, 1921) deserve a just appraisal. There is not any doubt that injection technique in embryological material is peculiarly susceptible to artefact and the meticulous presentations of Sabin (1902, 1911, 1913) seem almost impossibly fortunate.

If the venous origin of lymphatics is accepted, then their assistance in the return stream of fluid from the periphery is certainly in accord with behaviour which could be expected from modified veins. There is no doubt that the lymphatic endothelium acquires a morphological specificity very soon. In subsequent growth in the normal course of events, or in the phenomenon of injury requiring repair, the new lymphatics arise from pre-existing lymphatics by a building process. In this way they conform to the normal principle of homoplastic regeneration. Clark and Clark (1937), using the transparent chamber technique have established beyond doubt that new lymphatics do not arise de novo in the connective tissue in extra-uterine life in the rabbit.
THE LYMPHOID TISSUE

The definition of lymphoid tissue is not precise. Any tissue consisting predominantly of lymphocytes has been held to constitute it, Drinker and Yoffe (1941). The other components vary in character and in proportion. In all but the smallest aggregates of lymphocytes, connective tissue and reticulo-endothelial elements are present, and vascular tissue, either lymphatic or blood, is almost as consistent in its presence. In animals below the mammalia the tissue is characteristically more coherently integrated with the myeloid tissue; this association is less evident in the adult higher forms, but is a notable phenomenon in them in certain morbid states. Even in normal man the separation is not complete in an anatomical sense, since lymphoid tissue nodules are probably constantly present in the bone marrow, Dominici (1902). Later the relation of lymphocytes to myeloid cells will be discussed more fully, here it suffices to state that some evidence points to the existence of a much closer relationship than is generally accorded.

The definition of lymphoid tissue can be confined more closely without error as tissue consisting predominantly of lymphocytes together with their progenitor cells and other close relatives set in a vascular stroma. It has already been emphasized in Part 1 that in reticulo-endothelial organs the parenchyma arises from the stroma. This interpretation is a general one and applies more particularly to the cellular maintenance of the organ. In the first origin of lymphoid tissue it might be supposed that the stroma developed first but this has never /been
been confirmed, the origin of the first lymphocyte is still unsolved. It is akin to the famous riddle one encounters first in the nursery, 'which came first, the chicken or the egg?' The sophisticated reply, namely, 'the egg - but it was a reptilian egg', only half solves the difficulty but it supplies an analogy for comprehension of the primordial anlage of lymphoid tissue.

The term lymphoid tissue is preferred. It is older than other names suggested and it is very widely adopted. According to Erich, (1929,a) Aschoff maintained that the word lymphatic should relate to the tissue in question when germinal centres were present and the term lymphoid restricted to formations without them. It is doubtful whether anything useful is gained by this distinction, besides it reflects too static a concept of the morphology.

In the study of lymphoid structures it is immediately obvious that the appearances show topical variation. Deposits of the tissue may be circumscribed or diffuse, the cellularity of the formations may be high or low, in particular the density of lymphocytes may be uniform or the reverse. Recently it has become fashionable to draw a broad distinction between loose lymphatic tissue and dense lymphatic tissue but many special terms are used in addition. Generalisations about the lymphoid tissue are few because it is exceptionally difficult to study it comprehensively. It is exceedingly diffuse and widespread so that total ablation experiments cannot be applied with precision.

Aschoff considered that the lymphoid tissue organ could be regarded as distributed in three main groups.
1. The tissue in the lymph nodes. This part has afferent and efferent lymphatic vessels related to it and lies chiefly in the lymph stream.

2. That in the mucous membranes. Efferent lymphatics only are present and the tissue is located in the fluid streams going from the mucosae into the interior of the organism.

3. That in the spleen, where neither efferent nor afferent lymphatic vessels exist but the tissue lies in the blood-stream.

Quantitatively the bulk of the lymphoid tissue is accounted for in this preliminary schema, but not all of it. The facts are that at present there is insufficient knowledge and agreement for any more comprehensive account. An attempt is made below to describe the various moieties briefly.

While the general principles of this classification are adhered to, some tentative inclusions will be added.

At the outset several general remarks are offered. Lymphatics and lymphoid tissue are present in all vertebrates. They are first recognised in the Selachii (Elasmobranch fishes). It is probable that the fundamental reasons for the system lie in the attainment of increasingly better blood and blood vascular apparatuses. The spleen and the organ of Leydig are the oldest representatives of the actual lymphoid organs and in this primitive animal the spleen occupies its original site in very close proximity to the bowel. This has been interpreted as reflecting the old association of phagocytes with the coelome, and is a significant phylogenetic principle. Lymphoid tissue seems destined to
form round tubes, not only the alimentary tube, but later round other tubular structures in which there is flow. As the phylogenetic scale is ascended migration of lymphoid tissue from the old para-enteronic sites commences and reaches its highest expression in mammals, where, although the bulk of the tissue is still probably in the primitive position (disregarding the peculiar migration of the spleen), much is cut towards the periphery in the form of lymph nodes. The old lymphoid organ — the paleo-lymphocytogeion and the new, the neo-lymphocytogeion are relatively distinct in many ways and these include properties of interest to the pathologist. It is a general principle that disease has most chances of success in attacking more recently developed structures and functions; the old ones are more immune. This is particularly well illustrated in certain acquired diseases of the central nervous system, e.g. dementia paralytica. Examples can be cited for the lymphoid organs. Exclusive or almost exclusive involvement of the para-enteronic lymphoid tissue in primary tumour is exceedingly rare. The pseudoleukaemia gastrointestinalis of Briquet (1838) is so little known that under ten cases have been described (Symmers). (The original case is depicted in Cruveillier's Atlas (1835-42). By contrast Hodgkin's disease without extensive lymph node lesions is extremely rare, and intestinal or gastric lesions are recorded in under 5 per cent of cases in most series. Further, Hodgkin's disease is the commonest primary growth in lymphoid tissue and is characteristically a lymph node affection.

The break up of large lymphoid tissue masses into smaller separate parts is generally held to be an indication of phylogenetic maturity. 

/Thus
Thus it is thought that the dispersal of the pancreas of Asellius into several hundred mesenteric lymph nodes is a higher development.

The schema adopted to describe the lymphoid tissue is as follows.

1. The lymph nodes.

2. The sub-epithelial lymphoid apparatus which includes:
   (a) i. The lymphoid tissue related to the intestinal tract.
        ii. The lymphoid tissue related to the respiratory tract.
   (b) The lymphoid tissue of the integument.

3. The isolated follicles.

4. The thymus gland.

5. The spleen and haemal lymph nodes.

6. The ectopic lymphoid tissue.

1. The lymph nodes.

   The term lymph node is replacing the older one, lymph gland. This is an advantage because true glands are typically epithelial structures and characteristically they elaborate secretions. Lymph nodes are not epithelial and do not secrete in the ordinary sense of the term. From its derivation the word 'node' can be accepted as referring to a lymphatic channel knot or morass of Gordian complexity. (The French ganglion lymphatique conveys the same idea). Incidentally the word gland means an acorn and was originally adopted to describe the glans penis. Lymph nodes seldom resemble acorns; if they must be characterised generally, they are bean-shaped. These structures are situated in the lymph stream; as a corollary it is reasonable to suppose that they bring about modifications in the lymph itself and this is being demonstrated with increasing conviction.
In quintessence they are situated in the path of the circulating lymph. Topographically they are mostly organised in groups which are related to the larger blood vessels. It appears that afferent lymph is required to traverse a node after it has completed part of its journey to the blood. Large aggregates of nodes are found at the base of the neck, the roots of the limbs and the lymphatic hila of viscera or groups of viscera. Peripheral to these sites formed lymph nodes are rare and where present they are small. The actual disposition and identification of the nodes or groups of nodes present much difficulty. Despite statements to the contrary, it is certain that individual variation is considerable, indeed capricious. The meticulous demonstrations pictured by Poirier and Charpy, Cuneo and Marceille, and Jamieson and Dobson in Gray's Anatomy must be either reflective of individual cases or refer to an ideal. The number of lymph nodes in man is doubtful. Sappey (1869) hazarded that between 600 and 700 might be recovered by ordinary dissection but that many small ones might be overlooked. Drinker and Yoffe (1941) suggest the true number is probably nearer 2000.

2. (a) i. The para-enteronic lymphoid tissue is developed in relation with the epithelial lining of the alimentary tract. It is represented by the pharyngeal tonsils of Luschka, the palatine tonsils, the Peyer's patches and the less coherent or solitary lymphoid follicles of the entire tract. It is believed that these deposits are with few exceptions related to the lymphatic system solely by efferent channels. The generalisation is derived chiefly from studies of small representative parts.
parts. The solitary follicles may be relatively closely grouped as in the tongue, or dispersed as in the stomach and the colon. The aggregate of the latter is often impressively shewn in enteric fever and may be demonstrated in the normal by clearing the bowel in lactophenol. Orsków (1901) using this method computed their number to be between 2000 and 3000.

2. (a) ii. The lung parenchyma is normally devoid of lymphatics and lymphoid tissue but both are present in the immediate vicinity of the bronchial tree. The para-bronchial lymphoid tissue is generally diffuse and scanty but comparatively large, well-formed intra-pulmonary lymph nodes may occur. Embryologically the lungs are formed by outgrowth from the primitive pharynx and accordingly this component of the lymphatic system is virtually para-enteronic. An elaborate lymphatic plexus is situated in the visceral pleurae and occasionally small laminate lymph nodes are encountered here.

2. (b) The lymphoid tissue of the integument is probably considerable but very few studies have been made on it. Among the better known representatives are the solitary follicles of the conjunctiva (Bruch's glands). The problem of differentiating 'chronic inflammatory foci' from normal lymphoid formations is frequently difficult but it is possible that many foci of this description in the skin are in fact natural. Their relation to the lymphatics is unknown.

3. The isolated follicles. Very small deposits of lymphoid tissue are regularly encountered in the bone marrow, the uterine endometrium, the salivary glands, and the breast. The significance of these islets
is obscure. It has been suggested by Jordan and others that the bone marrow lymphoid tissue gives rise to the blood cells but this view does not yet enjoy much popularity. The recent work of Yoffee promises interest in this subject. The endometrial follicles may possibly have led to the opinion formerly widely held that the endometrial stroma was of lymphoid character.

4. The thymus gland is sometimes considered along with the epithelio-lymphoid apparatus. There is not complete agreement however that the tissue is truly lymphoid. For long the early development of the gland was disputed but it is now widely agreed that an entodermal outgrowth forms the anlage. Whether a proportion of cells become physiologically equivalent to reticulum cells or whether the mesenchyme cells at the locus do this is not settled though the latter interpretation is better supported. Morphologically the thymocytes have not any specificity by which they can be differentiated from lymphocytes, their mutual identity is probable. Serologically they are the same cells and the thymocyte’s high sensitivity to x-rays is exactly parallel to that of ordinary lymphocytes. The gland does not appear related to lymphatics in any particular fashion and lymphatic sinus tissue is absent from it. (Lymphatics are present in the stroma trabeculae and drain to the deep cervical and mediastinal nodes). This anomalous and peculiar structure is probably the most bewildering and labile lymphoid organ. Among the few facts known about it, one of the most striking is its dramatically rapid shrinkage in response to infection, a process which is almost akin to autolysis. Atrophy and necrosis sometimes subserve physiological
needs, a familiar instance is the universal death of the superficial part of the skin, which phenomenon enhances the insulative properties of the integument enormously. It is attractive to speculate whether or not acute thymic involution has some equally valuable function but the issue is an extremely complex one.

5. The spleen and haemal lymph nodes are unique representatives of the lymphoid tissue. The former was originally a haemopoietic organ and even in man an ephemeral phase of this activity is shewn in foetal life. Other functions have been superimposed but it cannot be said that any of them appear very vital. This lymphoid tissue is virtually in the bloodstream and constitutes the largest single deposit in man. Although developmentally para-enteronic it migrates peripherally and characteristically exhibits a tendency to react along with the lymph nodes in disease. The comparative anatomy of the organ is peculiarly inconsistent, at least two types of spleen are described in the mammalia. In the less common one, represented broadly by the spleen of ungulates the intravascular position is much less definite. It is tempting to wonder if this viscus is a kind of unfinished experiment in morphology and function.

The existence of true haemal lymph nodes in man is not universally acknowledged. Originally described by Gibbes in 1884 they have evoked desultory attention. The original description is poor but later publications have remedied this deficiency: Warthin (1901) gave an excellent account. They are small, inconspicuous, and usually about as difficult to find as the organ of Zuckerkandl or the superior parathyroid glands. The classical site to attempt their recovery is in the /inconsiderable
inconsiderable areolar tissue between the renal artery and vein. It is improbable that they are of much importance and personally I am only doubtfully convinced of their specificity; erythrocytes accumulate so readily in ordinary lymph node sinuses as the result of minimal trauma.

6. **Ectopic lymphoid tissue** is a phrase coined to describe deposits of the tissue in sites where it is not generally recorded. This phenomenon will be more fully discussed later, it includes problems related to the neo-formation of lymphoid tissue, including lymph nodes: and the significance of the phenomenon.

**The biological significance of lymphoid tissue**

The essential component of the tissue is the mass of lymphocytes. These cells are among the most mysterious in the body, singularly little is really known about them. Drinker (1946) emphasizing this, stated that the two acknowledged facts of importance regarding them are their high sensitiveness to x-rays and their indispensibility in the life process. The first characteristic is common to all lymphocytes in whatever species they may be found. The anatomical diffuseness of lymphoid tissue militates against the successful resection of it in entirety, although the skilful and ingenious operations of Sanders and Florey (1940) go close to realising it. The method of irradiation has been suggested as an alternative; but experimentation by this device has not been successful since the dosage required is lethal. Jolly (1924), reported that this sensitiveness could be considerably depressed if the irradiated lymphoid tissue were simultaneously deprived of its blood supply, which suggests that the susceptibility was not intrinsic.
It is a general principle that an actively proliferating tissue is more vulnerable to x-rays than an indolent one and a reasonable corollary derived from the lymphocytes behaviour is that it must be an actively growing tissue. The postulate is however difficult to reconcile with the apparent inertia so commonly observed in ordinary histological preparations of lymphoid tissue. Evidence has been adduced that the lymphocyte population of the blood vascular system is renewed daily or even several times a day. These results are derived from counts made of lymphocytes in the thoracic duct lymph. This at once raises the question - where do they go? A satisfactory answer is not yet available. It has been suggested that some are returned to the lymphoid tissue, others to the bone marrow, many to the intestinal lumen (yet they still disappear as readily in an animal previously deprived of its gut!), some to the skin to form epidermal epithelium, and some to the tissue spaces. This last site seems probable since lymphatic endothelium appears to resent the re-entry of lymphocytes. The idea that they are destroyed in the blood stream now receives less support.

The indispensibility of the lymphocytes is emphatic. It has long been suspected that the lymphoid tissue is part of the body's defence mechanism. This appears to have originated from the supposed efficacy of the filtering property lymph nodes exhibit. This effect is well shewn in nodes draining tattoo marks, or anthracotic lungs. The filter is not perfect because distant lymph nodes may be pigmented and even the spleen may become anthracotic. When the particles are really small e.g. bacteria or viruses the filtration is far from perfect. It is
established however that lymph nodes are important sites of antibody formation though the roles of the various cells concerned is unsettled; Oakley, Warrack, and Batty (1949). Ingenious interpretations have been offered e.g. the macrophage component elaborates the antibody which it then sheds in surface films, this material, claimed to be \( \gamma \) globulin, then becomes the cytoplasm of lymphocytes which is identified as \( \gamma \) globulin too, and these on their dissolution yield this fraction to the blood plasma. Other theories hold the lymphocytes to be directly concerned with the whole process and more recently the plasma cell has come into favour. This problem has prompted a great deal of work in the last few years and it is still too early to assess it.

Many clinical facts seem inconsistent with the too ready acceptance of an effectual defensive role on the part of lymphoid tissue. Enlarged tonsils and adenoids are not convincingly associated with high resistance to infection, nor is the florid para-enteronic lymphoid tissue of the child related to singular freedom from bowel infections.

The meaning of lymphocytic infiltration round cancerous growths is quite obscure. The phenomenon is very common but not invariable. I have to record one curious exception. In the series which I have studied of mucoid cancer deposits in organs or lymphoid tissue the phenomenon has been usually strikingly absent. In one relatively early metastasis in a lymph node the mucoid material was seen lying in the marginal sinus and in the vicinity a process of lymphocyte disappearance was evident.

The elementary function of lymphoid tissue is sometimes stated to
be the production of lymphocytes. Superficially this might appear axiomatic but it is exceedingly hard to prove. In morphological studies with fixed human tissue the best reasons for believing it are that they do not appear to be produced elsewhere, and adhering to the principle that cells are best recognised by the company they keep.

Since the primary interest in this work as a whole is Hodgkin's disease, emphasis will be laid on the more directly relevant problem of lymphoid tissue which the disease raises.
The structure of lymph nodes with observations on physiological human nodes

Appreciation of morbid lymph node histology presupposes full knowledge of the normal structure. The acquisition of this axiomatic requisite is difficult because good descriptions are few, and the latitude of normal appearances presented by lymph nodes is wide. In many standard anatomical texts illustrations of several easily procured mammalian nodes have long been offered as adequate. His (1860), a pioneer of descriptive microscopical anatomy based his structural schema solely on the lymph nodes of the ox, and certain of his drawings were repeated at intervals in other books for about fifty years. His described the lymph node as composed of a peripheral cortex of dense lymphoid tissue enclosing a sinusoïdal medulla. Von Recklinghausen (1871) first disputed the correctness of this and over the next thirty years better descriptions became available. Especially valuable were the works of Ranvier, Jolly, Flemming, Dominici, Gulland and Bunting. At the same time deficiencies still attended the descriptions; species differences were commonly overlooked and generalisations persisted without criticism. There has been relatively little published which applies exclusively to human lymph nodes until comparatively recently. Hellman (1913) and Heudorfer (1921) published very good accounts and illustrations of human nodes but the texts are not very accessible. In English, Bremner and Weatherford (1948) also used human material but the section is brief. A further anomaly was justly criticised by Job (1922-3).
who pointed out that the structure given in most text-books related to an ideal with little indication that all nodes were not the same.

The conventional description of a lymph node includes the following features. The node is invested by a capsule which is usually invaginated at one or more points to form a hilar intrusion; trabeculae spring from the inner aspect of this envelope. It is emphasized that with the possible exception of the Australian opossum the trabeculae do not characteristically segment the node. (In the Eutheria, the pig is often erroneously credited with the possession of septate nodes). In the hilar intrusion the major blood vessels arrive and leave and the efferent lymphatics accompany them. The multiple afferent lymphatics enter at any point of the capsule piercing it directly or after running a short distance obliquely. Under the capsule is the marginal or peripheral sinus. This is a sub-capsular lake from which the trajectorial sinuses which penetrate the node substance are given off. Subsequently these drain via the medullary sinuses into the efferent channel. The lymphatic tissue is disposed as pulp and follicular tissue. Some authorities refer to the pulp as medulla, an unfortunate convention since it is not anatomically the medulla, other terms are the parenchyma, or loose lymphatic tissue.

In the anatomical medulla follicular tissue is much less in evidence and the sinuses are separated by columns of loose lymphatic tissue which are described as the lymphatic cords. The sinus tissue is usually clear of lymphocytes, except in the trajectorial paths, it is lined and fenestrated by modified reticulum cells variously termed littoral cells
or endothelial cells. The former term is supplanting the latter.

Before describing the individual tissues listed above a general statement of node structure is appropriate.

Fundamentally a lymph node is a circumscribed plasmodium of reticulum confined and partly segmented by condensations of the fibrous elements of the surrounding areolar tissue. This locally condensed vascular connective tissue constitutes the capsule and trabeculae, the latter are generally very imperfect in man. The reticulum symplasma shews local attenuations and condensations and corresponding with these loci morphological modification of the reticulum cells is seen. An argyrophile filigree of reticulin is intimately related to this plasmodium and it too exhibits local alterations including focal deficiencies. In most of this basic framework are inserted lymphocytes and their precursors, together with a variable but small content of other cells.

This statement appears at first to have little relation to what is seen in an ordinary haematoxylin and eosin stained paraffin section. None the less it is correct, but suitable material and suitable methods are required to rationalise the identity. The general reticulum of nodes is most convincingly demonstrated by the old device of pencilling. A young lymph node is obtained as fresh as possible and without prior fixation is sliced, and macerated in Ranvier's alcool-en-tiers (33 per cent spirit), for 24-48 hours. Frozen sections are then cut in gum arabic. These are placed in the dilute spirit in a watch glass over a dark background and gently tapped with a moist soft camel's hair brush. As this is done a faintly opalescent fluid comes away with the medium,
this is a suspension of the mechanically dislodged cells, chiefly lymphocytes. The finished preparation is mounted in glycerin. Perfect results are difficult to obtain, but in parts of most specimens the continuity of the reticulum of sinuses, follicles and pulp is well shewn. In the centres of the follicles the reticulum is extremely delicate and this portion is almost always damaged or lost. The argyrophile reticulin filigree is demonstrated by silver impregnation. The method of Mitchell and W. Locki, (1944), gives extremely good results but is expensive. The reticulin mesh is closer in the anatomical cortex of the node than in the medulla and this pattern persists in old age (Denz, 1947). It is customary to counterstain these preparations with diffuse stains e.g. saffranin, neutral red, & etc, but these are substituted with light haematoxylin with advantage. The mesh is deficient in the centres of follicles. It is to be noted that the walls of small blood vessels are also outlined by these silver methods and further it is nearly always possible to trace optical continuity of the reticulin with the collagen of trabeculae which stain various tints of brown. Mallory's phosphotungstic acid haematoxylin, Lieb's haematoxylin, van Gieson's stain, and picro-fuchsin will also display some of the reticulin mesh in most cases. These methods supplemented by serial sections prepared by orthodox histological methods enable a satisfactory understanding of lymph node structure to be gained.

The Capsule and Trabeculae

Nodes are invariably set in areolar tissue and it is usual to
encounter anchoring strands of connective tissue running between this adjacent areolar tissue and the capsule. The capsule is conveniently regarded as a condensation of the peri-nodal connective tissue and not as intrinsic to the node. This interpretation is prompted by consideration of the embryonic development. In human nodes the envelope is composed almost entirely of white fibrous tissue which fibrifies progressively as it matures but rarely is the process complete. It is generally four to ten cells thick but variations with local thickening, extreme attenuation or even partial disappearance are seen, especially in old nodes. In its substance run blood vessels, lymphatics and less constantly nervous structures. Small islets of adipose tissue and lymphoid tissue are also sometimes encountered. The smooth muscle of blood vessels and elastic fibres from the same source are the only contributions of these elements in human nodes. Nerve fibres are fairly common in node capsules but are not apparently distributed in the node itself, paccinian corpuscles and ganglia are found rarely. The trabeculae are derived from the hilar intrusion and from the general capsule. In form they are very irregularly fenestrated laminae. This can best be appreciated from the study of serial sections and from corrosion specimens. The basal area from which they spring is very irregular in extent.

The blood supply is distributed from the hilar intrusion and trabeculae. These septa act as scaffoldings for the vessels in their earlier course. In the medullary tissue of nodes vessels may be seen running almost free of support in the lumina of lymph sinuses, having
left their perivascular connective tissue sleeves. Small branches leave these stems and enter the node pulp. A complex pre-capillary and venule mesh is formed by the terminal vessels. Around follicles, either of resting or pale centre type, an anastomotic basket-work is common, and a central arteriole generally supplies the centres of follicles. The blood vascularity of most nodes is seen to be remarkably high if measures are taken to ensure its visibility. The simplest method is auto-injection. As soon after death as possible the cadaver is placed so that passive hypostasis engorges the nodes it is desired to study. Within a few hours, when the subject comes to necropsy the node vessels are often satisfactory filled with blood.

The sinus tissue

After traversing the capsule the afferent lymphatics open into the subcapsular sinus. (This term is more descriptive than the alternatives peripheral or marginal since these suggest it exists only in one place). This space is rarely so complete as conventional description indicates; in many old nodes the underlying lymphoid tissue is in direct contact with the inner aspect of the capsule. The subcapsular sinus is continuous with sinuses which surround the fibrous trabeculae and these following a centripetal course traverse the anatomical cortex. The actual path taken may be tortuous and this is commonly due to displacement by the follicular tissue. In addition to juxta-trabecular sinuses others make their way independently in similar fashion. In the anatomical medulla the sinus system is usually more conspicuous because the lymphoid density is less. The efferent lymphatics effect junction
with these medullary channels and often also with the subcapsular sinus, and leave in the hilar intrusion. The sinus system is best seen in nodes related to the abdominal viscera especially the stomach, because in these glands the cellular structure is looser and this often is further enhanced by oedema, and physiological activity. The sinuses are lined by modified reticulum cells (specialised endothelial cells, littoral cells), and traversed by very similar cells arranged in retiform fashion. It is emphasized that these cells are but slight local modifications of reticulum cells. Downey (1922), in a very exhaustive paper demonstrated this beyond all doubt, and abrogated the mistaken view that they are true lymphatic endothelial cells. Macrophages, lymphocytes, plasma cells, granulocytes, and erythrocytes are commonly present in small numbers.

Reticulin strands continue across the sinuses in addition to forming a fenestrated wall. In developing nodes and in old inactive ones these fibres frequently appear naked, and constitute most of the internal structure of the sinuses.

Studies of morphology alone have limitations but it is probable that sinus tissue has not the permanence which might be imagined; it is further likely that sinus tissue can develop locally in response to appropriate stimuli. Conway (1937) led evidence that the sites of follicles were not constant, and while this is not universally accepted e.g. Denz (1947), it is a reasonable belief and reinforces this idea.

The lymphoid tissue

This exists in two primary forms, the pulp or loose lymphatic
tissue, and the follicles. In adult human lymph nodes the former may be the sole expression, at least at the level studied.

The pulp is composed of the general reticulum in the interstices of which are inserted lymphocytes and other cells. This free cellular component exhibits considerable variation in its composition dependent on the activity of the node. In a quiescent or inactive node the cells are nearly all mature small lymphocytes; whereas in the reactive state their proportion falls due to the appearance of lymphocyte precursors, proliferating endothelial cells of Marchand, plasma cells, and cells in mitosis, with occasional addition of granulocytes, giant-cells, & etc.

This groundwork lymphoid tissue displays peculiar focal aggregates of cells at intervals in its existence. These are termed lymphoid follicles, or less accurately lymph follicles. Unfortunately the different varieties have a confused nomenclature, and for so small structures they have roused much controversy. Two varieties are well recognised and can be demonstrated easily in human material. The first is the solid or resting follicle. These consist of closely packed small lymphocytes which form aggregates of elastic size. They are by far the commoner formation visible in human lymphoid tissue, being present throughout life, even long before birth. The conglomeration of small lymphocytes decreases in density peripherally and merges insensibly with the loose lymphatic tissue. The outline of these formations is conventionally circular in section but ovoid or bilobed configuration is common. Careful scrutiny reveals minute blood vessels and scanty reticulum cell nuclei in the lymphocyte mass. They are not traversed by /sinus
sinus tissue as a rule. Mitotic figures and pyknotic nuclei can generally be identified if sought. In common with most cells in mitosis their identity is usually a matter for speculation. The pyknotic fragments often present a bi-valve appearance which is peculiar to lymphocytes. In a silver impregnation deficiency in the reticulin fibres is generally present centrally, and the surrounding mesh is typically ragged in demarcation. These follicles are also termed secondary nodules because at one time the phrase primary nodule was used to indicate the segmented division of the lymphoid tissue produced by the capsular trabeculae. In recent works they are often termed the solid secondary follicles of Groll and Krampf.

The second conspicuous follicle is one in which there is a circumscribed spheroidal centre of large leptochromatic nuclei with symplasmic cytoplasm. The relative pallor of this formation in stained sections has led to it being termed a pale centre. This name, sanctioned by usage is not very imaginative, but carries less implication than others. The demarcation of this pale centre varies from remarkably sharp to indistinct. It is surrounded by a marginal zone of small lymphocytes which may encircle it uniformly or with increased density locally. In the latter event the maximum density is typically cap-shaped and orientated towards the capsule. These follicles have a long and interesting history in microscopical anatomy.

They were first described in detail by Flemming in 1885. He characterised them as possessing 'light centres with a dark shell in which the reticulum is often arranged concentrically'. He termed them /secondary.
secondary nodules in a morphological sense and physiologically 'germinal centres' or places of origin of lymphocytes. He pointed out that locally the pale centre was a site where mitotic figures were more numerous than elsewhere; that the displacement of the previously evenly distributed reticulum indicated an outward growth of cells from the centre and surmised that the daughter cells were driven out by a slow centrifugal pressure through the spaces of the reticulum. In addition he drew attention to minute stainable bodies of uncertain nature in between or in the cells of the pale centre. For some obscure reason nearly everyone prefers to call these by their German name viz. 'tingible korper'. In this country Bunting (1904, 1905), in his excellent account of the histology of lymph nodes added further details. He recorded that germinal centres were very rare or absent in the new born, and sometimes absent in adults. He noted their relative profusion at the periphery of nodes and paucity in the deeper parts. He observed too, that in active nodes the capsule might be mammilated by florid growths of nodules of this variety. In pencilled sections he observed that the germinal centres tended to drop out and adduced evidence of an entirely protoplasmic reticulum in the pale centre. Among the cells of the pale centres he recognised two main varieties. One had a large oval pale staining vesicular nucleus, and the other was smaller and darker. Occasional phagocytic cells, 'angular reticulum cells', were also present and he emphasized the very dubious cytoplasmic demarcation of all cells in the centre. He supported Flemming's contention that the centres were lymphocytopoietic by primitive micro-dissection methods. While this view is probably still the most widely supported by authoritative opinion
opinion today, the sceptics began to question it soon. Ehrich (1929) recorded that the first healthy doubt was that of Marchand in 1913, and in that year and the following, Hellman refuted their lymphopoietic properties with some very convincing facts. This last writer, who has considerable claim for attention, observed that the marginal zone of small lymphocytes did not appear to augment or decrease in regular fashion with coincident activity of the pale centre; that there was no definite relationship between the number of mitoses and the size of the marginal zone, and that convincing transition form between the centre cells and small lymphocytes simply did not exist.

Further to these he added the fact that typical germinal centres did not appear in the young until several months after birth. (In a later publication he modified this after finding that they sometimes arose in cases of foetal infection). His final reason was and is overwhelmingly convincing to a morbid anatomist, namely pale centres are not found in lymphatic leukaemia. In his later work Hellman, with Heidelberg and others introduced the idea that the pale centres were 'reaction centres' related to irritation.

Latta (1921) studying the development of the intestinal tonsils in lepus discovered that the first free cells were mainly small lymphocytes which arose by differentiation of fixed mesenchymal cells, and in a well balanced argument decided that the so-called germinal centre was not a centre of proliferation of lymphocytes. He was among the first to submit that the 'tingible korper' of Flemming were nuclear remnants of degenerating lymphocytes. In the same year Nakahara and Murphy (1921) attempted
attempted to solve the problem by ingenious experimental methods. They found that after a brief exposure to dry heat there followed a sharp initial fall of circulating lymphocytes which was closely followed by a rise, often up to 200/300 per cent above the level in the intact animal. Study of the lymph nodes and spleen at this point revealed numerous dead cells in all parts of the lymphoid tissue except the pale centres. By 48 hours numerous mitoses appeared in the latter and this activity persisted. They also found that if mice were injected with an emulsion of homologous living tissue ten days before inoculation with a cancer graft that they were rendered relatively resistant thereby and shewed a fall in the number of takes. This immunity was associated with a distinct lymphocyte rise in the blood and the level often rose again when the tumour inoculum was introduced. Parallel with these phenomena two related phases of mitotic activity in the pale centres were demonstrated. In further experiments using very small doses of x-rays similar changes were produced. They concluded that these findings were evidence of lymphocytopoietic function of the pale centres; it is convincing up to a point, but the crucial question of how the pale centre mitoses account for the lymphocyte rise was not met.

Jolly (1922) accepted Flemming's hypothesis without demur, and added some interesting matter. After a formal description of the chief cells of the pale centre he stated 'beside these clear nuclei are found scanty small lymphocytes and cells which are larger with basophile and scanty protoplasm which resemble the germinative lymphoid cells of the embryo and the myeloblasts of the bone marrow. They are of the same nature and
here are called lymphoblasts or leucoblasts'. A foot-note explains that 'these are the same cells as cells called large lymphocytes in many recent works - this just leads to confusion'. (This reference is probably to Maximow). He proceeded - 'Some authors have advanced the view that the mitoses of the clear centres belong only to the conjunctive framework' (Ribbert in Ziegler's Beiträge, VI. V889), but Jolly insisted that mitoses could be seen in manifestly lymphoid cells.

A reconciliation between germinal centre and reaction centre is accepted by some authorities, indeed this is endorsed by Aschoff. West (1924) believed that the pale centres were lymphocytopoietic and at the same time centres of cell destruction.

Maximow's work is very well known and need not be repeated. A frequently quoted summary is available in Cowdray's text-book of Special Cytology (1932). Allowing for the nomenclature anomalies the relation of small lymphocytes to certain cells of the pale centres is parallel to that given by Jolly.

This work carried out over many years has met with fairly general approval. It is however worthy of mention that cat and rabbit material were used in preference to human in which Maximow himself agreed it was difficult to identify the cells named. One interesting finding of Maximow is that the centre can be traced through a cycle of different recurring phases.

Two further authorities deserve mention. Ehrich (1929 a,b) clearly supported Hellman's idea. In addition he described a third variety of lymphoid nodule as well as recognising and characterising transition stages.
stages between the solid and pale centre types. This new formation he called a pseudo-secondary nodule and he outlined a tentative schema indicating the relations between the different types:

```
Solid secondary nodules
↓
Flemming's secondary nodules
↓
Transition forms
? → Pseudo-secondary nodules
↓
Lymphoid tissue
```

with the appearance of the proliferation endothelial cell of Marchand, and plasma cells.

The pseudo-secondary nodule is a large formation, of the order of 3 mm diameter, with indistinct outlines and best appreciated with very low-power examination.

The importance of the proliferating endothelial cells of Marchand in lymphocytopoiesis was stressed. These cells are well characterised and their recognition is easy. Ehrich is probably correct regarding their nature viz. precursors of lymphocytes, but I have not been able to convince myself of the entrance of lymphocytes into the peculiar veins which he described. Drinker and Yoffe (1941) in their highly critical study incline towards the orthodox view but readily admit the inconsistencies inherent in it.

The foregoing summary is necessarily eclectic, the subject has an enormous literature. In the main my studies have been limited to morphology and I shall not attempt to survey the immunological and irradiation methods which have been employed to solve the problem beyond stating that I have not encountered any convincing proof that the pale
centres in man are producers of lymphocytes.

**Involutionary changes in human lymph nodes.**

In the practical study of adult human lymph nodes it is remarkably common to find evidence of atrophy. Cowdray (1942) has drawn attention to the fact that lymphoid tissue shews a disproportionate rate of ageing.

It is established that in man lymph nodes reach their most florid development in late childhood or in early adolescence. Thereafter they are prone to undergo atrophy. This phenomenon is seen earlier and more consistently in the peripheral nodes. A detailed study of the problem, covering some 300 nodes from the axillae and mesenteries has been performed by me for another purpose and here only a brief account of the findings will be given.

There are two common expressions of atrophy in lymph nodes. The first which is most characteristically seen in para-enteronic nodes is a progressive depletion of lymphocytes. This process is apparently the result of a failure of supply, and it progresses slowly, rarely is the de-population anything like complete. Simultaneous with the fall of lymphocytes the proportion of reticulum cell nuclei appears to rise and often sinus tissue is thrown into prominence. Follicles with pale centres are rarely seen in this form of atrophy and often secondary follicles are absent. In most case there is also regression of the connective tissue of the node and the trabeculae become fragmented or tenuous.

The other form which is well seen in axillary nodes is best described as fat replacement. Fat cells replace the lymphoid tissue in
two main ways. They may either appear as a diffuse irregular sprinkling of individual fat cells, or by the development of a conical process of adipose tissue which originates near or in the hilar cleft and excavates the node. This fat cone may be formed by local coalescence of fat cells of the first type but is usually a separate formation. In many cases both mechanisms are visible, the individual fat cells lying ahead of the fat cone. The lymphoid tissue recedes absolutely passively in front of the advancing fat. Sinus tissue appears slightly more resistant and these structures are commonly out-flanked till the reticulin gives way and then they disappear. Eventually the capsule is reached, this membrane persists till all the lymphoid tissue and sinus tissue have disappeared, it then also succumbs. In general the node does not tend to gastrulate or collapse, the capsule retains its position till it is extinguished piecemeal. On the other hand the lymphoid tissue falls back onto capsule while it persists, thus conforming to the general shrinkage pattern described by Thompson (1942). Ultimately the node may disappear almost entirely except for a tiny residual crescent of surviving lymphoid tissue and capsule. There is evidence that the fat is a new product. It is paler than the adult yellow fat in the perinodal areolar tissue and often the cells are larger and more uniform in size, recalling the tissue of lipoma. In my studies I am led to the conclusion that the nutrition of the subject bears a close relation to the degree of the phenomenon and its recurrence. It has probably more influence than age alone. It occurs earlier and much more consistently in the obese, especially in women. The process can be almost complete
and yet the locus is still a potential lymph node. It is not uncommon to find carcinoma metastases in the relics.

Regeneration is commonly observed in response to appropriate stimuli. This is well shewn in the axillary nodes of obese middle aged women who develop mammary carcinoma. This repopulation is simply a reversal of the atrophic process and was admirably described by Reubens-Duval and Fage (1909).

Neither of these atrophic processes is inevitable. Even in elderly people, over 80 years of age, remarkably intact lymph nodes can sometimes be found. Follicles with pale centres may even be present.

Recognition of this atrophic process is important in the present work because one very remarkable and consistent finding is that it is very rare indeed in lymph nodes which are the seat of primary tumour.

Much less commonly atrophy with fibrosis is encountered. In cervical, axillary, and mediastinal nodes this is definitely rare, but minor degrees are more often seen in mesenteric, retroperitoneal, and groin nodes. This fibrosis is vascular in origin and is primarily a hyalinisation of blood vessels. Only rarely is it extreme. In groin nodes particularly, augmentation of the hilar intrusion connective tissue contributes. It is emphasized that this fibrosis does not arise from the general reticulum.

Fibrosis arising in the reticulum is common, but fortunately the unimportant forms have usually an obvious cause. It is very common in nodes invaded by carcinoma metastases and in scrofula, both conditions which are generally easy to recognise. Despite statements to the /contrary
contrary, fibrosis in sinus tissue or general node reticulum due to prolonged sinus catarrh or chronic non-specific inflammation is extremely rare. As a practical corollary the finding of reticulum fibrosis without obvious cause must be viewed with the greatest concern.
THE DEVELOPMENT OF LYMPH NODES

Lymph nodes do not appear until the lymphatic vascular system is tolerably complete. They are first observed in human embryos at or after the 40 mm. stage (10 weeks), Hellman (1931); Le Gros Clark (1945); Bremner and Weatherford (1948). The process of their development in man receives little attention in many standard works on embryology; Frazer (1931), does not offer information and Arey (1941) gives only a formal synoptic account. Gulland, (1894), studied the problem in considerable detail but his material included sheep, guinea pig, and rabbit tissues in addition to human specimens. Briefly the mode of origin which he traced was as follows. The first indication of node formation was the development of a freely anastomosing plexus of lymphatic at the locus. (It is interesting to note that in Gulland's drawings of this stage, the nuclei of the lymphatic endothelium are identical with the connective tissue (mesenchyme) nuclei in the vicinity). The anlage of the node consisted of an ill-defined islet of condensed connective tissue surrounded by these lymph channels. 'Leucocytes' then appeared between the condensed connective tissue cells; these were not described but it appears likely that lymphocytes were meant, since their origin was ascribed to the thymus. While this mode of population is doubtful in the light of more recent work, it was a convenient and safe shelving of the problem since the thymus appears first in the 8 mm. stage (5 weeks). In a human foetus of 3 inches (75 mm, about 3½ months) the nodes still retained a primitive appearance, there were not any hila, the marginal sinus was not highly organised, and deep sinuses could not be recognised. Noteworthy changes at this stage included increased cellularity of the
node and the appearances approximated to the structure of a lymphoid follicle. Further illustrations were given by reference to preparations from foetuses of 90 mm. and 115 mm. In the former trabeculae were seen in the process of formation. The smaller trabeculae arise from the slips of intervening general connective tissue between the individual lymphatics forming the marginal sinus; the larger were attributed to the folding of the capsule due to unequal growth. In the latter specimen the new definitive blood supply, an arteriole, was depicted entering in the substance of a large trabecula. The node capsule, and this deserves special emphasis, was merely a condensation of the surrounding connective tissue. This important fact aids materially in the proper appreciation of capsular changes in morbid lymph nodes. The capsule is a pressure condensation product of the environmental connective tissue and not an integral part of the node.

The development of the deep sinuses of lymph nodes is only briefly treated in most texts. Downey (1922), in a most comprehensive paper, described their origin in the lymph nodes of the pig. The first developments were spaces in the general reticulum of the node, as these increased in number and size the lining reticulum cells took on endothelial characters. The original non-endothelial character is very well indicated in Downey's drawing No. 6 where comparison with two normal blood vessels in the field emphasizes the point. Connection of these developing deep sinuses with the lymph channels of the plexus in which the node is developing is secondary. A corollary of much importance derives from this interpretation, namely lymphoid tissue has vaso-
formative ability. Koschowitz, (1950), has arrived at the same conclusion by further morphology studies in relation to chronic inflammation.

In my own studies I have relied on human material collected personally from post-mortem and surgical specimens. These sources have provided embryos of 22 mm, 46 mm, 65 mm, 78 mm, and foetuses of 94 mm, and 210 mm. In addition specimens were obtained from still-births and infants dying in the neonatal period, some being premature. The youngest specimen (22 mm. embryo) was poorly preserved, but the remainder were moderately good. So far as the limited material permits the sequence of development follows Gulland's thesis reasonably closely.

In the most mature specimens in the group the appearances shew little mutual difference. Between the neonatal period and the end of the first year is a span I have been unable to study adequately, due to paucity of material. The main interest in this phase concerns the appearance of germinal centres. The general consensus of experience indicates that these are very rare before six months of age. With the exception of isolated examples, the childhood and adolescent epochs were also not represented in the series collected.

It will be appreciated that this mode of formation of the lymph node is unduly formalised. When serial sections are studied it is at once obvious that the appearance of a ring of contiguous or reasonably adjacent lymphatic channels is only manifest at certain levels of the node anlage. It is indeed doubtful whether this 'ring' is essential. It would appear that when the process is resolved to its simplest generalisation, the fundamental fact is that lymphoid tissue develops at /loci
loci immediately adjacent to lymphatics. Acceptance of this principle
carries implications which will be exploited later but I believe it to
be correct for the following reasons. In embryonic material lymph
node development is seen without the orthodox plexus formation; it
tends to occur in the axil of a lymphatic vessel dichotomy or where
nearby blood vessels, capillaries as a rule, form an analogous enclosure
with a lymphatic channel. This mode of origin was depicted by Klein in
Quain's Anatomy, (1912 Edn.) The development of most of the para-
enteronic lymphoid tissue is evolved without any afferent channels
assisting the process, and in adult life it is quite common to find
lymphoid tissue developed in the immediate vicinity of lymph nodes,
either in or just outwith the capsule. This phenomenon is common in
nodes which are functionally active, hyperplastic, or undergoing
reactive regenerative change. Recognition of this process is afforded
little attention, Jolly (1922) is among the few who appreciate it.

It is not certainly known when lymph node production ceases.
Gulland surmised that there were three orders of lymph nodes which
developed successively, and invoked the last generation to account for
apparent new formation of nodes in adult life. Jolly (1923) expressed
a similar conclusion. Neo-formation of lymph nodes is not generally
 accorded much recognition but has been reported by Bayer (1885) and
MacCallum (1928). The former writer maintained that nodes could arise
de novo in adipose tissue, but as MacCallum pointed out this is unlikely.
MacCallum suggested that they might develop in relation to pre-existing
lymphatics. It will be appreciated that this is a difficult subject to
/investigate
investigate but in personal observations reasonably convincing evidence
for MacCallum's theory has been found.

Morphological studies indicate that the first development is a
focal lymphocytic infiltration which forms a circumscribed nodule. The
cells at this early stage are almost all small lymphocytes. Reticulum
cell nuclei then appear in this aggregate and pale centre tissue is
subsequently developed. The size of the aggregate appears to determine
these progressive changes. The site is usually very close to blood and
lymphatic vessels and the latter becomes gradually more intimately
related to the lymphoid mass and ultimately forms the marginal sinus.
The propriety of regarding this new formation as a lymph node depends on
the fact that carcinoma metastases may occur in it. (In the
illustrations of this present work there is a record of these findings).

An indirect method of supporting the fact of neogenesis is to
enumerate the nodes in presumed normals and compare them with cases
where they are increased. This I have been able to perform only once
and while in this case an apparent two-fold increase was demonstrated,
the method carries less conviction.
THE NUMBERS OF THE LYMPH NODES

Estimates of the total numbers in the body have already been indicated. The establishment of this figure is a very formidable task, at the same time it would be extremely interesting to know it. The significance which attaches to this enumeration is considerable, it might lead not only to a far better appreciation of the biological import of lymphoid tissue throughout life but might also furnish a means of determining whether neo-genesis of lymph nodes occurs.

While it has been impracticable to attempt this full scale enumeration, it was considered that even on a restricted regional basis some absolute figure might be useful. This section records an attempt to establish the number of lymph nodes recoverable from the human axilla throughout life. The dissections on which the findings are based are tedious performances and the total number is small, however they give some indication of the probable ranges to be found.

The axilla was chosen as suitable for the purpose. This region has the advantages of being readily accessible to ordinary post-mortem dissection where mutilation must be avoided, and of being tolerably circumscribed anatomically. In addition the nodes are less superficial and exposed to trauma than the inguinal groups. The left side was selected empirically.

Identification of the Axillary Lymph nodes

Johnston (Gray's Anatomy (1935)) gives the following information. The nodes are relatively large and vary in number from twenty to thirty. They may be divided into five groups which are not sharply demarcated
from each other.

**Schema, after Gray**

I. A lateral group of from four to six nodes lies medial to and behind the axillary vein.

II. An anterior or pectoral group of four or five nodes lies along the lower border of the m. pectoralis minor in relation with the lateral thoracic vessels.

III. A posterior or subcapsular group of six or seven is placed along the lower margin of the posterior wall of the axilla in the course of the subcapsular vessels.

IV. A central group of three or four large nodes is imbedded in the fat near the base of the axilla.

V. An apical group of six to twelve nodes is situated partly posterior to the upper portion of the m. pectoralis major and partly above the upper border of this muscle and extends upwards into the apex of the axilla along the medial side of the axillary vein.

This schema is not easy to follow in the actual process of dissection. In the practical study the text of Taylor and Nathanson (1942) proved more valuable. These writers have considered the recovery of lymph nodes as a practical surgical problem and two features of their anatomical studies deserve particular mention. The first is the use of veins as indicators of lymph node locations. The veins are analogous to the axis of advance used by a division in battle, the nodes being related paravenously. Maintaining the military simile, just as
the component brigades and supports may be disposed in line, column or echelon, in similar fashion the nodes along tributary veins are conveniently described. The second advantage is a simplification of nomenclature which is really an outcome of the first idea. The subgroups are named according to the vein to which they are immediately related.

**Schema, after Taylor and Nathanson (1942)**

1. The axillary vein group, a variable but considerable group, they extend along the vein from the outer border of the m. pectoralis major to the apex.

2. The subcapsular vein group, a few lymph nodes lying on the axillary surface of the vein, the teres and m. latissimus dorsi; they are usually related to the subcapsular nerve as well.

3. The lateral thoracic vein group, a chain of nodes lying approximately in the mid-axillary line in close relation to the digitations of m. serratus anticus, often between the bellies from about the fifth rib up to the apex. They are related to the lateral thoracic nerve of Bell.

4. The thoracic-acromial vein group, the highest group in the axilla, essentially coinciding with the axillary vein group. They coincide with the entrance of the cephalic vein into the axillary vein.

In Gray's system the numbers given yield a minimum of 23 and a maximum of 34. Taylor and Nathanson do not submit figures. It will be seen that the two systems are almost synoptic the groups corresponding...
as follows.

<table>
<thead>
<tr>
<th>Gray</th>
<th>Taylor &amp; Nathanson</th>
</tr>
</thead>
<tbody>
<tr>
<td>Part of IV + I corresponds to II</td>
<td></td>
</tr>
<tr>
<td>Part of IV + III</td>
<td></td>
</tr>
<tr>
<td>IV</td>
<td></td>
</tr>
<tr>
<td>V</td>
<td></td>
</tr>
</tbody>
</table>

The contents of the left axilla were recovered from twenty routine necropsies. The patients' ages varied from 1 day to 81 years, eleven were females and nine were males. All cases were free of primary lymph node disease and did not suffer from cancer or scrofula. The methods adopted were simple. The axilla was rendered accessible with the widest reflection permissible and the arm extended at right angles to the body. The entire axillary fat was then dissected out. In the final clearing the m. pectoralis minor was reflected and a segment of the axillary vein removed. The larger nodes were readily appreciable with the bare fingers and easily picked out. This form of blunt dissection proved the best particularly where advanced degrees of fat replacement obtained. In these cases the slightly enhanced tenseness of a node and its just appreciable relative pallor proved valuable. Smaller nodes were also recoverable by palpation. The smallest nodes presented more difficulty. In cases where the fascia was not heavily fat laden stretching it often brought them into view. Where much fat was present they were best identified by compressing the teased specimen between two sheets of thick perspex and examining it against a strong light. Many of these nodes are very small, of the order, 1.0 mm. diameter. Finally
histological confirmation was obtained by making a single composite block of pieces from the presumed nodes recovered.

**Results**

<table>
<thead>
<tr>
<th>No. of Case</th>
<th>Sex</th>
<th>Age</th>
<th>Lymph Nodes recovered by dissection</th>
</tr>
</thead>
<tbody>
<tr>
<td>1.</td>
<td>F</td>
<td>1 day</td>
<td>11</td>
</tr>
<tr>
<td>2.</td>
<td>M</td>
<td>6 days</td>
<td>10</td>
</tr>
<tr>
<td>3.</td>
<td>M</td>
<td>30 days</td>
<td>23</td>
</tr>
<tr>
<td>4.</td>
<td>M</td>
<td>2 years</td>
<td>31</td>
</tr>
<tr>
<td>5.</td>
<td>M</td>
<td>5 years</td>
<td>32</td>
</tr>
<tr>
<td>6.</td>
<td>F</td>
<td>13 years</td>
<td>31</td>
</tr>
<tr>
<td>7.</td>
<td>M</td>
<td>18 years</td>
<td>30</td>
</tr>
<tr>
<td>8.</td>
<td>F</td>
<td>22 years</td>
<td>26</td>
</tr>
<tr>
<td>9.</td>
<td>M</td>
<td>29 years</td>
<td>29</td>
</tr>
<tr>
<td>10.</td>
<td>F</td>
<td>43 years</td>
<td>21</td>
</tr>
<tr>
<td>11.</td>
<td>F</td>
<td>47 years</td>
<td>24</td>
</tr>
<tr>
<td>12.</td>
<td>M</td>
<td>51 years</td>
<td>20</td>
</tr>
<tr>
<td>13.</td>
<td>F</td>
<td>56 years</td>
<td>23</td>
</tr>
<tr>
<td>14.</td>
<td>F</td>
<td>61 years</td>
<td>12</td>
</tr>
<tr>
<td>15.</td>
<td>M</td>
<td>65 years</td>
<td>15</td>
</tr>
<tr>
<td>16.</td>
<td>F</td>
<td>66 years</td>
<td>5</td>
</tr>
<tr>
<td>17.</td>
<td>F</td>
<td>69 years</td>
<td>9</td>
</tr>
<tr>
<td>18.</td>
<td>F</td>
<td>70 years</td>
<td>11</td>
</tr>
<tr>
<td>19.</td>
<td>M</td>
<td>72 years</td>
<td>8</td>
</tr>
<tr>
<td>20.</td>
<td>F</td>
<td>81 years</td>
<td>12</td>
</tr>
</tbody>
</table>

/Summary
Summary of Findings

The series recorded extend over the life span but the representation of successive periods is not uniform. This is primarily due to lack of suitable material; specimens from the first three decades were difficult to obtain.

In so far as the findings permit analysis the following observations are offered. At birth and in the early neonatal period the total number of nodes is in the region of 12. This low figure is attributable chiefly to the smallness of the structures, even at one month their size is much larger making their identification easier. Throughout childhood till the end of the third decade figures near the classical maximum are found. Thereafter there appears to be a steady slow decline. The figures are too small to assess sexual differences; it is my impression however that in the elderly, women have fewer nodes and this appears related to obesity which is commoner. Adiposity is apparently conducive to fat atrophy of lymph nodes (q.v.) besides which it renders surviving nodes very inconspicuous. Perfect illustration of the results would require photographs of all the nodes from each case, this was too expensive and only half the cases have been so pictured.
PART III

HODGKIN'S DISEASE

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BIOGRAPHICAL NOTE ON THOMAS HODGKIN, M.D. (EDIN.)

Thomas Hodgkin was born at Pentonville, London, on August 17th, 1798. His father John Hodgkin, to whom he was much attached, was a member of the Society of Friends and a paedogogue by profession. Thomas, who was the third of his four sons, was a premature child and was never robust in health. He was educated at home by his father who was no mean scholar, and acquired an enviable knowledge of Latin and Greek. Concerning his medical training little is recorded. Part of his course was performed at Guy's Hospital, but he also studied abroad, in Paris. He graduated M.D. from Edinburgh in 1823, a matter for delectation. His thesis was entitled 'De absorbendi functione' and he received special commendation for his Latin—a graceful recognition for an Englishman—from a Scottish University. In 1825 he joined the staff of Guy's Hospital, undertaking the offices of Curator of the Museum and Lecturer on Morbid Anatomy. These he performed with distinction for a period of twelve years, at which point a senior post of physician became vacant at that institution. Rather unexpectedly he was passed over and one of his colleagues, Dr. Babbington, obtained the situation. This circumstance was a considerable disappointment to him and he left the hospital. Some years later, in 1842, he was invited to undertake the task of reorganising the medical School of St. Thomas's Hospital in which he was appointed Lecturer on the Practice of Physic. Significantly enough, this hospital had previously been linked with Guy's—in fact Hodgkin's first appointment there was a consequence of this disruption. A lesser man might readily have exploited this state of affairs to cause...
an unpleasant rivalry but Hodgkin was not so minded.

On the formation of the University of London he was elected to the Senatus Academicus and proved a proficient and wise counsellor.

As a man Hodgkin was a Christian gentleman and one of the Salt of the Earth. Like his father he was a sincere member of the Society of Friends and adopted their dress and opinions throughout his life. He was one of those precious individuals who cannot remain apathetic observers of injustice and inhumanity: and it seems that much of his spare time was spent in advocating and supporting reforms. His work in this direction was essentially philanthropic in character and not political as some critics have suggested. Indeed, it is interesting in these days to note that he foresaw and warned against the tyrannies unprincipled trade unionism would exert on individual liberty. Altruistic principles seldom exist in those with worldly wisdom and Hodgkin had only the former.

His medical works include his Catalogue of the Guy's Hospital Museum, (1829); his paper entitled, 'On some morbid appearances of the absorbent glands and spleen', (1832); 'Lectures on the means of promoting and preserving the health', (1835); and his Lectures on the Morbid Anatomy of the Serous and Mucous Membranes, (1836–40). More is related concerning the first two entries elsewhere. His lectures on the health are dignified and sound. They must represent fuller expositions than the original verbal communications made at the Mechanics' Institute, since the book which contains three lectures – with appended notes, takes several days to read! The last publication /unhappily
unhappily was never finished; it was used as a standard text for some years and is superior to contemporary British works. It is really a general morbid anatomy text book because the subjects of the title are amplified considerably by appended observations and notes. It is not easy to read, but I was brought up to believe that anything worth while was invariably hard. The English translation of Rokitansky's Morbid Anatomy (1846) is far worse. His writings have been criticised as uninspiring and too diffuse, Hale-White (1944). The former charge is possibly true where the reader is a pharmacologist, but they do not leave a morbid anatomist unmoved. The second objection may be tenable in the sense that they are not written in the stilted and abbreviated scientific jargon of today. Theirs is the diffuseness of scholarship and there is in them a refreshing clarity of expression. The strong discipline of the Humanities of Hodgkin's early schooling is reflected in them. This renders them a trifle pedantic perhaps - even unconsciously humorous at times, but I for one am delighted to meet such old friends as the English equivalents of ut, quin, quominus and causa!

Particulars of Hodgkin's private life are anecdotal, one fragment which delights me personally is that at one point he owned a velocipede - a modern equivalent of which vehicle I myself possess. It is recorded that he was wont to travel from Lewes to Brighton on this machine.

He married comparatively late in his life, in 1850. There were no children of this union. He died from an attack of dysentery in Jaffa,
Jaffa, Palestine, in April 1866 while accompanying his friend Sir Moses Montefiori on an expedition to relieve the plight of destitute Jews. This gentleman erected an obelisk over his burial place - 'Humani nihil a se alienum putabat'.
THE CONTRIBUTION OF THOMAS HODGKIN

It is almost certain that priority must be conceded to Hodgkin for the initial characterisation of the disease which now bears his name.

In the introductory apology for his original observations he modestly disclaimed this honour. Indeed he suggested that the morbid alterations he was about to describe were already recognised and as likely to have been succinctly summarized. This circumstance has prompted several writers to sponsor the claims of previous observers, and certain of these will be considered below.

At the time of his now famous publication he was Morbid Anatomist at Guy's Hospital. This paper was read before the Medico-Chirurgical Society of London in two parts; the first on 10th January 1832, and the second a fortnight later. Hodgkin himself was not a member of this society and the communication was made verbally by his colleague Dr. Lee who was the Secretary. Despite certain limitations, especially the absence of immediate microscopical study this is a work of conspicuous merit. It has pleased several to emphasize its deficiencies and even to argue that Hodgkin is lucky to have been remembered at all. The blunt statement that Hodgkin 'undoubtedly had no conception that in one, or possibly two of his cases he was dealing with a peculiar and rare disease', Reed, (1902), and the patronizing account of Hale-White, (1924), are both unjust.

The study in question is based on the findings in six personally observed cases and a seventh reported from St. Louis Hospital, Paris, by M. Lugol. (In addition a further series of seven cases not relating to
The identities of the cases, after Hodgkin.

I. Joseph Sinnott, a child about 9 years old, in Lazarus' ward under Mr. J. Morgan. Dated 2nd November, 1826.

II. Ellenborough King, aged 10 years, in Luke's ward under Dr. Richard Bright. Dated 24th September, 1828.

III. William Burrows, aged about 30 years, in Naaman's ward under Mr. J. Morgan. Dated 28th November, 1829.

IV. Thomas Westcott, aged apparently about 50 years, in Clinical ward under Dr. Addison. Dated 8th January, 1830.

V. 'A middle aged man who had latterly been a patient of Dr. Back' previously under Dr. Bright. Not dated.

VI. Thomas Black aged about 50 years, in Barnabas' ward. Under Dr. Bright. Not dated.

VII. (Reported case of a man who died in the St. Louis Hospital, Paris in April (?1831).

In the interests of brevity, ease of comparison, and ready recognition of the salient facts it is expedient to present these findings in a more formal manner than the original. The relevance of several of the cases has been seriously questioned on many occasions. The exercise of critical study is reasonable, the responsibility of rejection resting with the individual. Accordingly an eclectic series will be given which represent the examples which I believe are true cases of the disease.
Case I. Joseph Sinnott

History. He had been ill for about nine months, complaining of pain in the back which extended round to the abdomen. His brother, a bed-fellow, died of phthisis a few months previously. On admission he was found to have ascites and oedema of the prepuce and scrotum. (There is no account of the patient's death).

Sectio Cadaveris

Head. The arachnoid mater contained a few 'opaque' spots and there was serous effusion under it. The pia was very thin and pallid. The brain was soft and flabby; no local morbid change was present in it.

Chest. The Heart was natural.

Pleurae Rt. Numerous old and tough adhesions were found. There was evidence of recent pleurisy in addition. Lt. A very few adhesions were noted.

Lungs. In the right organ there was a doubtful tuberculous cicatrix at the apex. 'but the substance of both lungs was generally light and crepitant with a very few exceedingly small tubercles scattered through them' Lucosa of bronchi, engorged. The bronchial glands were greatly enlarged and indurated.

Abdomen. Peritoneum. A sero-purulent effusion was present. 'The viscera were universally overlaid with a very soft light yellow coagulum, too feeble to affect their union though evidently having a tendency to do so' The mucosa of the stomach was very pale. The contents of the intestine were unhealthy. The mesenteric glands were enlarged, several reaching the size of a pigeon's egg. They presented semi-cartilaginous hardness and were streaked with black matter. The Liver was normal on the whole but dissection revealed a few tubercles somewhat larger than peas. These were white, semi-cartilaginous and of uneven surface.

The Pancreas was firmer than normal.

The Spleen was large, and contained many tubercles.

The Kidneys, both were mottled with a light colour but free from induration.

Para-aortic Lymph nodes (he adds) 'A continuous chain of much enlarged indurated absorbent glands of a light colour accompanied the aorta throughout its course, they were closely adherent to the bodies of the vertebrae and extended along the iliac arteries too'.

Thoracic duct. Large: the coats were transparent and normal.
Case II. Ellenborough King

History. Admitted 6th August 1828. Patient was the youngest of six children, the others being alive and well. He was fit till about thirteen months ago when his strength, flesh, and healthy appearance began to fail. A tumour was noted in the left hypochondrium, corresponding to the spleen. The glandulae concatenae on the right side were also enlarged, they fluctuated somewhat in size, this was attributed to treatment while in hospital. His appetite was good and he had not any haemorrhagic tendency. He became dropsical before death.

Sectio Cadaveris

Head. Not opened.

Neck. 'The glands in the neck had assumed the form of large smooth ovoid masses connected merely by loose cellular membrane and minute vessels: when cut into they exhibited a firm cartilaginous structure of a light colour and very feeble vascularity; but with no appearance of softening or suppuration'.

Chest. The pericardium contained more fluid than normal, the sac itself was not diseased. The Heart was normal. Pleurae. Old adhesions were noted. The lungs were generally healthy. The bronchial and mediastinal glands presented the same enlargement and induration as those in the neck.

Abdomen. Peritoneum. Ascites was present. The stomach and intestines were normal. The mesenteric glands were mostly normal, some were slightly enlarged. The para—sacral glands were grossly enlarged, and so were the juxta splenic and those accompanying the iliac vessels.

The liver was natural.

The spleen was enlarged to at least four times its normal size; 'its surface mammilated and its structure sprinkled with tubercles' presenting the same structure as the enlarged glands'. 
Case IV. Thomas Westcott.

History. Admitted ten days previously with a conspicuous lymphadenopathy affecting all the accessible glands, especially those in the neck, axillae, and groins. They were moderately firm and many were as large as pigeon eggs. Abdominal distension was present. The functions of the brain had been somewhat disturbed and the vision of the left eye was imperfect. He died very suddenly.

Sectio Cadaveris (4½ hours post mortem)

External. The veins of the head and neck were turgid.

Head. The arachoid mater was remarkably thick and opaque. The brain itself was normal; the cerebellum was small. The right optic nerve was smaller than the left.

Neck. Cervical lymph nodes. These were greatly enlarged, in general the deepest being maximal in size. They were set in loose cellular tissue and when separated were smooth and white. In consistence they compared with a testicle. The cut surfaces were white and homogeneous, vascularity being low. Degenerative change was not apparent in them.

Chest. Pericardium. Natural. Heart. The organ was much hypertrophied and dilated. The cardiac muscle itself appeared healthy. The agonal thrombus was very poorly formed.

Pleurae. Normal, cavities free from effusion.

Lungs. Both were pale, crepitant and mildly emphysematous. The bronchi contained some thick mucus.

Mediastinal lymph nodes. Those along the subclavian arteries and about the roots of the bronchi were much enlarged.

Axillary lymph nodes. These were considerably enlarged and shewed precisely the same features as those in the neck.

Abdomen. The peritoneum was not remarkable.

The mucous membrane of the stomach and intestines shewed no pathological features of note.

The Liver was very large, pale and slightly granular.

The Spleen was very greatly enlarged, measuring approx. 9 x 5 inches and proportionately thick. 'Its colour was lighter and redder than is natural, and more firm and close'. The cut surface was speckled with innumerable little white nodules of irregular outline, they were not tubercles.

The Pancreas was large and pale but otherwise healthy.

The abdominal lymph nodes. Those at the lesser curvature of the stomach, several in Glisson's capsule, and the para-aortic groups were greatly enlarged. The mesenteric glands though enlarged, bore no comparison in size to the former specimens.

(Note. Hodgkin is emphatic on this point, the disparity was remarkable). The lymph glands of the groins were greatly enlarged and bore a close resemblance to the axillary groups.
Case VI. Thomas Black.

History. Admitted nineteen days previously. He was affected with large tuberose swellings of considerable firmness on both sides of the neck, in both axillae and both groins. He was pale, cachectic, and dysphoeic. The abdomen was greatly distended. The cervical tumours were of about two years duration.

Sectio Cadaveris

The head was not opened.

Neck. Cervical lymph nodes. (See note below).

Chest. Pericardium and heart. (No record).

Pleurae. There was evidence of recent inflammation. Serous effusion was present. Lungs. (No record).

Mediastinal lymph nodes. The glands accompanying the carotid, subclavian, and internal mammary arteries were greatly enlarged. The posterior group were less conspicuous, and the bronchial glands did not appear affected.

Abdomen. The peritoneal cavity contained copious yellow serum mixed with some flakes of lymph.

Stomach and intestines. (No record).

The Liver was small and pale with an irregular and uneven surface. Two or three white nodules which resembled 'fungoid tubercles' were visible at the surface.

The spleen was normal.

The pancreas was embedded in grossly enlarged lymph nodes but intact.

The kidneys were livid and congested.

(The cervical, axillary, para-aortic and inguinal lymph nodes were reported collectively) sic:

'The tumours in this case very nearly resembled each other in structure; there was a little difference in firmness. They varied in size from that of a horse-bean to that of a hen's egg. They were mutually adherent in light fashion in their respective groups. The capsules proper were inseparable from the gland substance. The cut surfaces were pale and slightly translucent, some were of semi-cartilaginous hardness'.

Brief notes on the cases excluded

Case III. William Burrows.

(The findings in this case, which were first reported by H. Peacock, Esq., will not be set out in detail. The patient was known to suffer from syphilis. The morbid anatomy corresponds with this disease and the lymph node changes, which were confined to the lower half of the body, did not resemble the other examples).

Case V. A middle aged man.

(The identity of this case is unknown, the history is vague and the necropsy report does not indicate sufficiently characteristic findings to warrant its inclusion. Careful reading of the text rouses an uneasy suspicion that the record was not made as soon as it might have been - ego quoque miles sum).

Case VII.

'Cancer cerebriformis of the lymphatic glands and spleen'. (This case did not come under personal scrutiny by Hodgkin and is therefore omitted).
The conclusions expressed by Hodgkin

Certain of these, which are quoted in full, not only reflect Hodgkin's shrewdness and acumen but shew an astonishing insight into the unique entity he had discovered.

He wrote 'It may be observed that notwithstanding some differences in structure, to be noticed hereafter, all these cases agree in the remarkable enlargement of the absorbent glands accompanying the larger arteries, namely the glandulae concatinatae in the neck, the axillary and inguinal glands and those accompanying the aorta in the thorax and abdomen, that as far as could be ascertained from observation or from what could be collected from the history of the cases the enlargement of the glands appeared to be a primitive affection of these bodies rather than the result of an irritation propagated to them from some ulcerated surface or other inflamed texture through the medium of their inferent vessels, and that although, in some instances the glands so enlarged may contain a little concrete inorganisable matter such as is known to result from what is called scrofulous inflammation it is obvious that this circumstance is not an essential character but rather an accidental concomitant to the idiopathic interstitial enlargement of the absorbent glandular structure throughout the body. That unless the word inflammation be allowed to have a more indefinite and loose meaning than is generally assigned to it this affection of the glands can scarcely be attributed to that cause since they are unattended with pain heat and other ordinary symptoms of inflammation and are not necessarily accompanied by any alteration in the cellular or other surrounding /structures,
structures, and do not shew any disposition to go on to the production of pus or any other acknowledged product of inflammation, except where, as in the cases above alluded to inflammation may have supervened as an accidental affection of the hypertrophied structure'.

'It appears in nearly all cases to consist of a pretty uniform texture throughout and thus rather to be the consequence of a general increase of every part of the gland than of a new structure developed within it, and pushing the original structure aside as when ordinary tuberculous matter is deposited in these bodies'.

Concerning the deposits in the spleen he added '... has been found more or less diseased and in some thickly pervaded with defined bodies of various sizes, in structure resembling that of the diseased glands. We might from this circumstance be induced to suspect that these bodies in the spleen, like the enlarged glands themselves are the result of a morbid enlargement of a pre-existing structure'.... (Here he quoted Malpighi (1660) who considered the acini or granulations in the spleen to be glands). ...'Hence we may conclude that if, as I conceive to be the case, there be a close connection between the derangement of the glands and that of the spleen, the latter is a posterior effect, and on this account may not always have been produced when that of the glands or some other disease carried off the patient'.

Summarizing, his conclusions were: that it was a primitive affection of the lymph glands, without any obvious primary inflammatory cause; that it could not properly be called inflammatory but seemed the consequence of a general increase of every part of the glands and that caseation in the affected nodes was due to secondary infection (scrofula).

/Finally
Finally he considered that the splenic changes were of the same nature and probably usually followed the gland changes.

It is submitted that most pathologists who believe that this disease is neoplastic or a primary reticulosis would have little difficulty in accepting these conclusions at the present day.

Several minor features are of interest. It is to be noted that all the cases were males, and two were children. Dropsy is recorded in five of the seven and there is occasional reference to black pigmentation in the affected glands. This last finding is rarely alluded to in works after 1900, it may refer to products of haemoglobin destruction.
Note on Critical Analyses of Hodgkin's Series

Several authorities have performed these and awarded various identifications. These are briefly summarized in the appended table.

**SUMMARY OF ANALYSES OF HODGKIN'S SERIES**

<table>
<thead>
<tr>
<th>CASE</th>
<th>WILKS 1865</th>
<th>REED 1902</th>
<th>SYMMERS 1924</th>
<th>HALE-WHITE 1924</th>
<th>FOX 1926</th>
<th>NAEGLI 1932</th>
</tr>
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<tr>
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<td>+</td>
<td>?</td>
<td>?</td>
<td>?</td>
<td>-</td>
<td>+ (Tuberculosis)</td>
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</tr>
<tr>
<td>III</td>
<td>- (Lues)</td>
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<td>?</td>
<td>?</td>
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<tr>
<td>VII</td>
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<td>?</td>
<td>?</td>
<td>+</td>
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</tbody>
</table>

**TOTAL** | 4 | 1 or possibly 2 | 2 | 4 | 3 | 'probably all' |

+ = Considered Hodgkin's disease.
- = Not considered Hodgkin's disease.
* = No comment.
? = No indication given.

Personally I am satisfied with the identity of cases I, II, IV, and VI, Wilks' analysis.
THE CONFIRMATION OF THE IDENTITY OF THE DISEASE

This is a fascinating chapter in the history of the malady. By a most fortunate set of circumstances some of the material from the original cases on which the paper was based has been preserved. The post-mortem histology has been merely delayed.

In most institutions post-mortem material does not command the priority accorded to the surgical specimen routine. A lapse of ninety-four years is nevertheless exceptional and it is of interest to trace how it came about.

It is remarkable how comparatively recently the subject of paleontology has been exploited. Even as late as the eighteenth century the recognition of *cerauniae* as primitive tools of Early Man was sporadic and hesitant.

In the middle of the ensuing century Boucher de Perthes gained acceptance of his claims and the study rapidly expanded. After this general sanction was won, attention was turned to physical anthropology and gradually the subject of paleopathology came into being. It is a neglected subject on the whole, particularly in this country, contributions to it being scanty and anecdotal. The Americans exploited it better, because, like archaeology it is an expensive form of enquiry. For obvious reasons much of it concerns persistent structures like the bones and teeth. Various studies have been made from time to time concerning these; Moodie, (1923), gives many references. Muskens, (1926), in his huge treatise on epilepsy depicted a series of trephined neolithic skulls and suggested that the apertures were surgically produced.
for the relief of that malady. In several specimens evidence is present which suggests survival of the operation, at least long enough for a suppurative osteitis to develop. Malunited fractures of long bones have often been described, and it has been claimed that syphilitic and tuberculous bone disease has been recognised in neolithic man, though this is not universally accepted.

The next logical refinement was the application of histological methods. This is of course possible only in comparatively recent material.

Oliver Wendell Holmes in one of his whimsical asides (1859) wrote — 'there were jars in rows where interesting cases outlived the grief of widows and heirs in alcoholic immortality — for your preparation jar is the true monumentum aere perennius'.

So far as I can discover the late Professor John Teacher of this University was the first to take advantage of this possibility — but it is not claimed that he did so as a result of reading 'Elsie Venner'! In 1894 Teacher became under-keeper of the anatomical and pathological department of the Hunterian Museum (William Hunter 1718-1783). Six years later he produced the valuable and complete catalogue (1900). In the case of the tumours, Teacher prepared sections and remarked how surprisingly successful this proved, despite the fact that the specimens had been preserved in spirit for 120-150 years. It is quite gratifying to realise that this work anticipates American labours in the same field by at least twenty years! Ruffer (1921) is generally credited as the first to attempt this: his studies were related primarily to mummified tissues from the Egyptian excavations. On the whole the experiment was
not successful, the majority of the tissues being too brittle, even after special treatment. Shaw (1938) reported the results of histological examination of the mummy of an Egyptian singer, Har-mose who lived in the 18th Dynasty (circa 1490 B.C.) The results in this case were exceptionally good.

In 1926 Fox examined several original specimens from Hodgkin's cases in Guy's Hospital Museum by modern histological methods. It is interesting to note in passing that Bright's disease as well as Hodgkin's has come under microscopical scrutiny, Osman (1937). It is greatly to be regretted that only three renal specimens of Bright have come down to us, all unhappily soundly injected with red lead. (One shewed changes of amyloid disease, another subacute extracapillary glomerulo-nephritis and the third chronic glomerulo-nephritis).

The identity of Hodgkin's material is not entirely straightforward. Since this is so fundamental an issue it will receive separate attention.

The source of the original material

The specimens which have been examined histologically by Fox (1926) and MacCallum (1928) were recovered from the museum at Guy's Hospital. This institution was founded by Thomas Guy in 1725 but for the greater part of a century it was under the aegis of St. Thomas's Hospital which also enjoyed a large share of the same patrons discreditably amassed fortune. Shortly before Hodgkin joined the staff, Guy's had become fully independent and its energetic secretary, Harrison, was launching forth schemes for its improvement. He purchased a collection of morbid anatomical specimens from a private source and this formed the nucleus
of a museum which Hodgkin rapidly augmented. Hodgkin related how in the space of four years the collection amounted to over three thousand, a great increase on the original five hundred. In 1829 he published his catalogue. Like all catalogues it represented an enormous amount of thankless work and considering it was the first proper inventory it is a very creditable achievement. It represents one of the best classification systems of its time, shewing a great advance on the famous museum at Leyden where only five headings were considered, viz. Ossa Morbosa, Partes molles morbosae, Calculi, Monstra, and Varia.

The entries vary much in completeness: some have clinical notes and full descriptions; others are very brief, e.g. 'another of the same'. In a quiet way it ranks high in entertainment value. It is delightful to know that Lt./Col. Herriot of the 22nd Regiment of Foot presented the museum with the skeleton of an Elephant! (943). Museums were museums in those days, not merely a set of teaching bottles! (Possibly it was the sort of gift that could not very well be refused).

Hodgkin used a simple consecutive number system to identify the specimens. This method has limitations. The most obvious difficulty being that the arrangement of cognate subjects in groups with serial numbers is impossible; the addition of fresh material introduces numbers not in the series already allotted. However, it is doubtful whether the adoption of modern library methods is much better because classifications themselves are continually liable to change. The important principle is that once any particular system is introduced it should not be altered - usually a pious hope.
The identification of Hodgkin's specimens

The sources of information available include Hodgkin's catalogue (1829), Wilks' papers of 1856 and 1865, and Fox's paper (1926).

The information yielded by a critical study of these may be summarized as follows.

1. **Hodgkin's Catalogue. 1829**

   There is direct evidence of specimens from Case I, Joseph Sinnot. The relevant entries are 1558 and 2009. On the negative side there could not be any record of specimens from Case IV, Thomas Westcott or Case VI, Thomas Black since both are dated after 1829, the date of this edition. Case V cannot be identified at all since there is not any name, number nor date. It is unlikely that any specimens from Case III, William Burrows would be recorded here since it is dated late 1829 (November 28th). Absence of information concerning Case II, Ellenborough King, is curious since necropsy was performed in the previous year, presumably the specimens were still being prepared.

2. **Samuel Wilks' paper of 1856**

   In this communication Wilks gave his cases his own serial numbers. Some of these are identified further, among them Cases 41, 42, and 44. From the text there is definite evidence of specimens being present in the museum at that date from Cases I. Joseph Sinnot (Wilks 41); II. Ellenborough King (Wilks 42) and IV. Thomas Westcott (Wilks 44).

3. **Samuel Wilks' paper of 1865.**

   In this paper there is evidence of specimens from five of Hodgkin's original series.

   /Case I.


Case IV. Thomas Westcott. Specimens 1538.50.; 1555.29.; and 1558.50.; All of glands.

Case VI. Thomas Black. Specimens 1543.32.; 1543.64.; and 1858.30. nature not stated.

4. Fox. (1926)

Fox was unable to identify the reference numbers given by Wilks 61 years previously, with specimens presently in the museum. He learned however from Mr. Burne, Curator of the Royal College of Surgeons and Dr. Beadle, Curator for Pathology, that one of Hodgkin's cases was in Guy's museum No. 1523 - under the name of Dr. Richard Bright. This statement is difficult to reconcile with Hodgkin's Catalogue entry No. 1523, which refers to a case of fungoid testis. The number must have been re-used, presumably between 1829 and 1856. This specimen, Fox declares, can be well identified with Hodgkin's Case II because its description coincides. An extract from the present (1926) catalogue identifies it as spleen and lymph nodes from Case II, Ellenborough King, and adds: 'See Insp. 6.P.156 and Prep.1541.12 Sec Edit' - this last number is the same as Wilks' 1865 reference to the cervical lymph nodes from the same case. Fox further learned that the present specimen 4768 corresponds to 1541.12.

He also obtained two further pieces from tissue studied by Hodgkin. These were present number 4769, identified with Case IV, Thomas Westcott, and present number 4770, identified with Case VI Thomas Black.
It is presumed that material from I, and III no longer exists.

The results of Histological examination

The spleen and lymph nodes from Case II Ellenborough King shewed a typical microscopical picture of Hodgkin's disease as presently defined and understood. The photograph made by Fox is unmistakable and convincing. Jackson and Parker, (1949), employed a closely similar photograph, probably from the same source, as a frontispiece to their book.

The preparations from Case IV, Thomas Westcott, were not so satisfactory but still strongly favoured the diagnosis. In Case VI, Thomas Black, Fox regarded the lesion as an example of 'The large lymphoid cell sarcoma type of growth' but he also declared that coarse fibrosis was present in the specimen. Eosinophiles are not mentioned in this case — perhaps it is too speculative, but might this not be an example of Warthin's type II Hodgkin's sarcoma?

MacCallum(1928), also confirmed the identity of the disease and stated that it shewed the characteristic morphology brilliantly.

This tangible evidence is curiously satisfying and vindicates the pioneer efforts of Thomas Hodgkin. At its lowest it fully meets the minimal assessment of Reed, (1902), at its highest it anticipates the kinship of lymphoid tissue sarcomata.
NOTE ON SUPPOSED REFERENCE TO THE DISEASE BY 
WRITERS PREVIOUS TO HODGKIN

It has long been the custom in medical writings to introduce a subject with germane historical facts. These may be a genuine history but often they are thinly disguised premisses for later argument, or provocative allusions to stimulate interest. Hodgkin's disease made its début so quietly that it was a comparatively long time before it too gained the distinction of a history. Now we have the authority of Holy Writ for the observations 'that of books there is no end', and 'Is there anything whereof it may be said - See, this is new, lo! - it hath been of old time which was before us'. Hodgkin, good Quaker that he was had these facts in mind even in the first sentence of his paper. This anticipation was apparently rapidly rewarded because he adds a footnote on page 97 sic. - 'shortly after the reading of this paper, I was favoured with the following communication from my friend G.O. Heming of Kentish Town:-

'Dear Sir,

You will, I am sure, be pleased with the following extract from Malpighi.

Yours truly,

G.O. HEMING'

'In homine difficilius emergunt (speaking of the granules in the spleen): si tamen ex morbo universum glandularum genus turget, manifestiores redduntur, aucta ipsarum magnitudine, ut in defuncta puella observavi, in qua lien globulis conspicuis racematis dispersis totus scatebat'.

//Hodgkin
(Hodgkin did not need to be told this, he was familiar with Malpighi's work and in fact alluded to this reference on p. 88). The exact context of this information is nearly always given wrongly, it appeared first in 'De Viscerum Structura exerçitatio anatomica', Chapter 5, page 124. These figures apply to the original edition printed at Bonn in 1666 — and not the London edition which appeared three years later. It is also available in Malpighi's 'Opera Omnia' which was printed in 1687 (Figuris elegantissimis in aes incisis illustrata!) in London. It appears on page 111 in Tom II of this edition.

Gowers (1879) stated that Malpighi was the first writer to mention this association of general enlargement of the lymphatic glands with nodules in the spleen. It is very questionable if this can be assumed to mean anything more than exactly what it said. The characterisation is far too incomplete to sustain the claim that this refers to Hodgkin's disease. Several other diseases could produce just such a picture. Writers previous to Hodgkin had alluded to, or described instances of general enlargement of the lymphatic glands, but most of these were regarded as scrofulous or carcinomatous. A remarkable example of the former is attributed to Morgagni/ (1752).

A claim is also submitted from time to time on behalf of Craigie. In the first editions of this authors 'Elements of Morbid Anatomy' (1828), under the heading of 'vascular sarcoma' or 'enlargement with induration' of the glands, is distinguished a form of lesion which could represent Hodgkin's disease. ...'The great hardness and the malignant tendency of this growth have procured for it from most authors the ominous names of /scirrhus
scirrhus and cancer. Though correct enough for all practical purposes these epithets are not justified by the anatomical characters'. (In the 2nd Edition of the work which appeared in 1848 there is not any significant addition to this statement). Fox (1926) interpreted this account to reflect recognition of a morbid process possessing features of both scrofula and neoplasm. Wilks (1856), though possibly unaware of Craigie's idea, restated the same reflections - as a dilemma. This particular book of Craigie's dealt with disease in a rather abstract fashion, actual cases were not cited to any extent. Had it been otherwise the sponsorship might well have been Scottish.
The Nomenclature of the Disease

If Hodgkin defaulted in this respect - the deficiency has been remedied by others. Wallhauser writing in 1933 collected fifty one synonymous titles which have been employed to connote it. Many are trivial variants, but several are still in current use. The eponymic Hodgkin's disease is very widely known, even if it is not universally used.

The French equivalent La Maladie de Hodgkin and the German, Die Hodgkinsche Krankheit, are less widely used than formerly. In English this old name has a stark arresting quality with a certain aura of mystery. It has the advantage of being non-committal, and has undoubted claim to priority.

A common alternative is Lymphadenoma. This is relatively old, having been suggested by Wunderlich in 1866. Etymologically it is doubtful and there is little to commend it. Some authors (Robb-Smith et al) sometimes add 'verum'. If this is necessary then the word must be condemned. In point of fact Wilks and Moxon (1875) defined lymphadenoma as 'any tumour having a lymph gland-like structure' but this has either been abandoned or forgotten.

Lymphogranuloma or lymphogranulomatosis enjoys a certain popularity among continental and American writers. It suggests that the disease is granulomatous in nature, a theory which is still not proven, and it is insufficiently specific. Malignant granuloma conveys an idea to which Ewing has contributed so much; this concept is discussed elsewhere but as a name it is a difficult hybrid which is liable to confuse those unacquainted
unacquainted with it. Malignant lymphoma, Bilroth, or the less antithetical lymphoblastoma of some American writers is too vague since many other growths could properly constitute it. The names suggested by those who regard the disease as a reticulosis have been considered already.
THE SUBSEQUENT HISTORY OF THE DISEASE

An enormous amount of work has been performed on the subject of Hodgkin's disease. It is impossible to give a comprehensive account here, and particularly in an outline, the just apportionment of credit to those who followed Hodgkin in the study of the malady is very difficult. It has already been noted that the disease has masqueraded under different names and this adds further obstacles in the path of historical research. Suffice it to say that the contributions which have accrued over a period of more than a century are overwhelmingly numerous. Popular interest in the subject has its crests and troughs as its nature, cause, and remedy are alternately discovered or discredited.

Many accounts of the main facts are available in text-books, reviews and papers. In examples of the foremost, in English, a very good summary is that of Gowers (1879), and another of merit is that by Murray (1909). More recent contributions of this character are those of Gordon, Gow and Rolleston (1937) and Jackson and Parker (1947).

Circumspect and unprovocative reviews have had a certain popularity, particularly in the United States of America, but to a lesser extent in this country. Widely quoted publications of this variety were offered by Longcope (1903); Rolleston (1925); Simmonds (1926); The Rose Research on Lymphadenoma (1932); and Wallhauser (1935). A lengthy review supported by nearly six hundred references was produced in 1948 by Hoster, Dratman, Craver, and Rolnick. This is possibly the most comprehensive of its kind as an amassment of information, but it is so

/non-committal
non-committal and uncritical that it loses much of its appeal.

Individual papers, particularly when they are the work of one man undoubtedly form the most stimulating source of information. This is because the writer is an individualist with a purpose, and he who goes alone goes furthest. The articles by the following are conspicuous because reasoned criticism is brought to bear on the historical aspects related. Wilks, (1856 and 1865); Reed (1902); Gibbons (1906); Ziegler (1911); Oliver (1913); Mueller (1921); Warthin (1930); Ginsberg (1934); Krumbhaar (1934); Symmers (1948); Custer and Bernhard (1948) and Jackson and Parker (1949). This by no means exhausts the list of valuable contributions and others will be referred to later, but as in most controversial subjects some dismally bad writings exist, inaccurate and stilted assemblies of second-hand material where even the references are unreliable.

In the necessarily discursive reading undertaken for the purposes of this work many little publicised points of information have emerged. Below I shall endeavour to offer an account of the title of this section in the form of an eclectic summary with criticisms. It is reasonable that I should disclose my bias since this will facilitate my later purposes but for this I am naturally personally responsible. I believe that the history of Hodgkin's disease is best presented in three parts which cover respectively the periods 1832-1865, 1866-1903, and 1904 - to date.

**Period I 1832-1865**

Up till the close of this period the history is anecdotal. The publication of its sponsor was manifestly not received with much /enthusiasm
enthusiasm. This was partly due to the fact that medical journals had a more restricted and smaller circulation than nowadays, and also because Hodgkin did not suggest a distinctive name for the disease. This trivial short-coming was in full accord with his lack of worldly wisdom but was a severe impediment to its advertisement. A name has remarkable power—witness the notoriety the 'Pancoast syndrome' has gained. Hodgkin's work excited scant recognition even from his own colleagues. There was however one heartening exception. Richard Bright, writing in 1838, gave full credit to Hodgkin; and judging from his remarks he appreciated the reality of this unique disease. The value of this acknowledgment is emphasized by reflection on the exceptional ability of Bright. This minor episode of loyalty is as solitary as the single company of infantry which Platea sent to Marathon and it seems to have suffered a similar fate.

It has already been suggested that tentative recognition of the disease anticipated Hodgkin's description. Malpighi (1666), and Morgagni (1752), possibly; and Craigie (1828) certainly knew of it. It seems improbable that it was a new disease like Trench Nephritis or von Economo's encephalitis lethargica. Although this postulate is speculative, it appears reasonable. The incidence of Hodgkin's disease is low but apparently fairly constant, and therefore it was liable to be recognised sooner or later by careful observers even in the absence of a conventional description. It seems very likely that Velpeau (1840–1) recognised it independently in France. This writer discussing the aetiology and pathology of lymphoid tumours in a lecture—demonstration/exposition
exposition clearly appreciated the existence of primary growths unrelated to infective disease or ordinary cancer. In the case described, a young woman, the indications are that it was the same morbid process.

In 1845 Craigie of Edinburgh described leukaemia and several months later, Virchow (1845) also did the same. It is probable that both writers knew of Hodgkin's disease but at this period it was still in the aleukaemic leukaemia group which had not been fully investigated.

In 1853 the disease reappeared in British literature. Markham described a case under the heading, 'Fibrinous deposits in the Spleen, thoracic glands & etc.' These specimens were removed from the body of a man aged 30 years. The naked eye morbid anatomy was well described, particularly that of the spleen which was of the hard-bake variety and weighed 1 lb. 10½ oz. (740 g.) This material was submitted to microscopical study by Bristowe at the same meeting. A fresh unstained specimen was examined and the report which stressed the great fibrosis present is strongly suggestive.

Three years later (1856) Wilks' first paper was published. This was entitled 'Cases of Lardaceous disease and some allied affections, with remarks'. This is a long article, of interest chiefly because of the evidence it furnishes concerning the preservation of material from Hodgkin's cases I, II, and IV. (Cases 39, 41, 42 and 44 of Wilks' series; case VI is reported). He described 'Hodgkin's disease' fairly well and certainly knew a good deal about it. As Hale-White (1924) insists, it stands greatly to Wilks' credit that he acknowledged Hodgkin's priority. (The cynic might point out that both Hodgkin and
A belated recognition of the disease was recorded by Pavy (1859), he wrote:— 'The malady is so striking and yet so peculiar that when carefully studied it is almost impossible to mistake its identity'. One case, occurring in a man aged 30 years was described briefly.

In 1865 Wilks' second paper appeared. This included a critical analysis of Hodgkin's cases with an assessment of their accuracy. He added other eleven but they are not all included in the text. A little microscopy was given in some of the cases, and one feels that he almost realised the features several times sic:— 'They were firm and composed of tough fibro-nucleated tissue' - 'composed of cells which somewhat resemble those of tubercle'. Unfortunately his discussion was vague but in summarising he observed shrewdly that the disease presented, 'some features of tubercle, some of cancer'. The hardships which Hodgkin strove so much to mitigate were still present, 'Case 12, Sarah P— aet.13. Under Dr. Gull, February 13, 1864. Her parents were dead, and she had consequently been very badly fed ...' An important immediate outcome of this paper was that the disease received its classical name.

Period II 1866-1903

In the first period it has been seen that after a somewhat precarious transit the disease has arrived at the stage of being an entity complete with a name and patron.

In this second phase the real trouble starts and yet it is from here, despite the confusion, that it will emerge in its full dignity and preposterous ambiguity. In the same year in which Wilks' second paper
appeared Cohnheim identified some cases of aleukaemic leukaemia with what he termed pseudo-leukaemia (pseudo-leukämie). This concept, or perhaps more accurately this name, has come in for much adverse criticism. The central idea however was excellent because it emphasized the unity of the primary morbid process — and literally it meant a leukaemia that played false. Since it is now widely appreciated that leukaemia is but an inconstant expression of certain reticulo-endothelial system diseases the idea is justified. Rather than despise it as a 'misch-masch' it is better to consider it as premature. The usage persisted especially in the German Schools, (Paltauf, Langhans, etc.). Cohnheim himself in 1870 identified 'certain cases' of Hodgkin's disease as pseudo-leukaemia and these doubtless corresponded to the 'hard' variety of Paltauf. In this year also, Cornil collected a series of cases of Hodgkin's disease, of which the description was now current, and added two more of his own with a careful study of the morbid anatomy. Trusseau (1868) introduced the term 'L'adenie' as a Gallic synonym for the malady. The word is variously rendered as 'adénie', 'adenia', etc. by other authors. His contribution was primarily clinical, the morbid anatomy was indifferently recorded.

After this point the main interest lies in the determination of the microscopical appearances by modern histological methods. Although histology was the earliest refinement of anatomy and pathology it was relatively elementary till this era. (Hassel's microscopical anatomy (1849), while praiseworthy to a degree, is comparatively elementary).

In 1872 Langhans described the results of microscopy in a case of pseudo-leukaemia; the account gave some details of the multinucleate /giant
giant cells and dense fibrous tissue in the lesion but it was brief and not illustrated. Six years later, in 1878, Greenfield of Edinburgh gave a further description. This certainly has priority, a point which is understandably emphasized by Alumni of the Capital (Muir 1924, van Rooyen 1938, and Ogilvie 1940), and remarkably enough by dwellers in Albania (now Londoniana), including Robb-Smith (1938). I have studied this paper carefully and frankly I feel it falls a little short of its reputation. It recounted precisely and carefully the clinical features of five instances of the disease. It is of interest that one of these, Case I, a man of 26 years, was under the care of Murchison who eight years previously had described the recurrent fever of Hodgkin's disease. (This original case was a child of six years who remained under observation for several months before death). Symmers (1945) is to be congratulated for taking the Profession to task for attributing this discovery to the Dutch and German physicians, Pel and Ebstein. The mode of preparation which Greenfield used is not described but it seems probable, judging by contemporary articles, that the sections were stained with logwood. The multinucleate cells were well described and line drawings depicted them; Greenfield considered they were particularly related to the trabeculae of the new fibrous tissue. He stressed fibrosis as integral to the complex - 'The growth has an essential tendency to induration' and he observed the frequency of perivascular fibrosis. Many critics have insisted that Greenfield maintained that increase of lymphocytes, a 'simple' lymphoid hyperplasia was one of the first evidences of the disease. There is not any direct statement to this effect in the text. A carefully worded account of the changes in adjacent
adjacent lymph nodes beginning to enlarge was given but the stress was unquestionably - and perfectly correctly too - on the incipient fibrosis. In the description of the splenic lesions the black pigment recorded by Hodgkin was also noted by Greenfield. He also observed that in the liver the disease began in the portal canals. Greenfield appeared to draw a distinction between Hodgkin's disease and lymphadenoma. (This interpretation is antithetical to that of Jackson and Parker (1949)). The latter term was widely employed in a generic fashion at this date although it had been suggested by Wunderlich as a specific term for the disease twenty years previously. Dr. F.E. Reynolds a former assistant with the late Professor Greenfield tells me that Greenfield was too good a pathologist to venture into print lightly, so it is probable that he said much less than he knew.

Doubtless as the result of limitations of staining methods, the eosinophile cells in the new tissue escaped notice till 1892, when they were discovered and fully depicted by Goldmann.

The remaining contributions of importance in this period are those of Sternberg (1898), Andrews (1902), Reed (1902) and Longcope (1903). These writers it will be noted belong to three different nationalities and anyone who reads much on the subject of Hodgkin's disease cannot fail to be impressed by the partisan tendencies of their respective compatriots. The complete lack of paeans for Andrews is to be regarded as a sort of negative self-effacement characteristic of the British. At the same time Andrews deserves the Palm for one of the most lucid and accurate descriptions of the microscopical features of the disease that has ever been written. His communication was /contributed
contributed to a symposium on the subject, called primarily to settle whether tuberculosis was an aetiological factor or not.

The paper was written in a refreshing personal style, a little repetitive perhaps, but evidencing a clear insight into the disease from his own personal experience. He commenced his study with a definition which is worth quoting - 'I mean by it a disease characterised by a progressive enlargement of the lymph glands and lymphoid tissue of certain internal organs, notably the spleen and liver; unattended by the indiscriminate metastases of sarcoma and characterised by what I believe to be a definite histological change in the lymphatic tissues affected'. Then followed an excellent description of a normal lymph node so arranged that the pathognomonic changes in the malady were most aptly compared and contrasted. Among the features described he emphasized the following:

The general plan of the structure of the node was obscured and simplified, the distinction between cortex and medulla being almost or entirely abolished so that an unwonted homogeneity resulted. Lymphocytes were reduced in number to an extent which rendered the framework of the node more readily visible. Corresponding with this there was hyperplasia of the framework and the fibrillar reticulum became conspicuous, this change amounting to actual fibrosis. The endothelial cells in connection with it were more numerous and more obvious. Scattered individual cells of this description attained a very large size with two, four, or more nuclei. These cells were especially numerous in the soft rapidly growing forms of the disease, and were quite different from Langhan's giant-cells. They most closely resembled
resembled the large cells found scattered throughout many rapidly growing sarcomata. Eosinophiles were sometimes but not always greatly increased. The affected nodes were singularly exempt from degenerative changes but these did occur occasionally.

In his ultimate summary he compared the lesions with tuberculosis, and, whilst inclining to the view that it was a granuloma insisted that this was in a confined sense, a point being reached when all similarity ceased. This account is necessarily abbreviated, there are many small points raised in the original text which revealed an appreciation of the peculiarities of the malady which was singularly penetrating.

Sternberg (1898) gave a very complete description of the pathological findings in fifteen cases, thirteen of which were considered to be in the category pseudo-leukaemia. The clinical histories are meagre in the extreme. The histological findings concurred mutually. In eight (i.e. out of thirteen) positive evidence of tuberculous disease was present in one or more organs, and Sternberg concluded that the disease was a peculiar form of tuberculosis. Necropsy was performed in only one case in the relevant series. (It may interest readers that this publication is exceedingly difficult to obtain. I am indebted to Mr. Archibald Goodall, F.R.C.S., F.R.F.P.S.G. for procuring a micro-film copy of it from U.S.A. The illustrations were extremely good).

In 1902 Reed published her paper. This work is very widely known and probably the most freely quoted in relation to the histological study of Hodgkin's disease. It is difficult to suppress one's feelings entirely and for this reason the severity of the authoress, and her irritating style detract from the undoubted merit of the work. It
must be stated that the description is very good, and the deductions, inferences, and speculations are sound. Several interesting new observations are added. These include, the apparent increase in vascularity in the nodes in the earliest stage, the occasional presence of Touton type giant-cells and the mode of fibrosis. In her series of eight cases three died and necropsies were performed. Reed discounted entirely the role of tuberculosis as an agent of the disease but believed that it was the commonest mode of death.

The publication with which this phase closes is that of Longcope. This is one of the most thorough expositions and is probably the best of its kind. It includes detailed protocols of his eight cases, experimental work, and so far as I am aware the first systematic examination of the bone marrow to ascertain whether the eosinophiles emanated from thence. In three of his cases out of four that came to necropsy he reported evidence of a response to the demand for eosinophile leucocytes but rightly cautioned against too sanguine an interpretation of findings in a tissue which is difficult to sample accurately. His efforts to transfer the growth to monkeys failed completely. He presented a careful and thoughtful summary emphasizing the anomalies inherent in accepting the disease as either inflammatory or neoplastic but was unable to solve the dilemma. It is interesting to learn from this author that Langhans, (1872), was the first to conceive of the extra-nodal deposits of Hodgkin's disease as being autochthonous in character. This is an unexpected priority - antedating Dominici (1900) by almost thirty years.

/Period III
Period III 1904 to date.

With the close of the second period there followed a dissemination of the characterisation of the disease which enabled others to assist in the researches which it prompted. Almost every aspect has now been examined with competence and thoroughness. It is an unhappy truth that the last fifty years of study have contributed refinement but only limited progress. While some discoveries have been added, others of previous time have been forgotten. Much of the knowledge gained is negative, but this is not equivalent to useless.

In this section, the subsequent history will be traced on a restricted basis with particular attention to problems which most directly concern the pathologist. These include contributions to the morbid anatomy, attempts to elucidate the nature of the disease, and classifications of the disease with corollaries developing from them.

**Advances in Morbid Anatomy**

Additions to the morbid anatomy are few in this period. In English the most important are the studies furnished by Symmers, (1924), and Pullinger, (1932).

Symmers' work was a very thorough examination of 14 necropsy cases with interesting and thoughtful interpretations of the findings. Among the points brought out was emphasis on how the true lymph nodes bore the brunt of the disease in contradistinction to the submucosal lymphoid tissue which generally escaped. Symmers envisaged an 'auxiliary lymphoid tissue' which normally was inconspicuous in amount, but which augmented greatly in Hodgkin's disease and frequently provided new loci in which the lesion might develop. He was among the first to consider
Some of these may well have been chronic myelodendritic leukemia superimposed on myeloid metaplasia or myelofibrosis.
that while an enlarged cervical lymph node might be the first clinical sign of the disease, this often betokened established lesions in the deep mediastinal nodes. The analysis of the findings in various sites including the spleen and liver, conformed to the general experience. His speculations and arguments on the nature of the disease were reasonable and it is interesting to record that he anticipated Medlar, (1931), in his recognition of the morphological similarity between the marrow in chronic myeloid leukaemia and the cellular complex of Hodgkin's disease. Since he regarded the fibrosis component in terms of residual inflammatory scarring the absence of collagen from the marrow picture did not impress him. (His two photographs of these lesions admittedly shew convincing similarity, but single high power fields are not a conclusive comparison).

Pullinger's contribution to the Rose Research on Lymphadenoma, (1932), comprised an exceptionally complete examination of the microscopical features of the disease together with observations on atypical examples. The evolution of the lesion from its incipient stages to maturity was traced and described in detail. Pullinger followed the topical development of eosinophiles from lymphocytes in the lesions using Mann's stain. This metamorphosis she considered as indicative of a myeloid affinity in the new tissue. She reinforced this thesis by drawing attention to the almost invariable presence of polymorphonuclear leucocytes in addition to eosinophiles. She was the first to emphasize this feature and also described how the neutrophiles were often visible as small coherent aggregates. Pullinger's study is undoubtedly the
most meticulous work which has been offered on the microscopy of this disease. The illustrations are copious and extremely good, and the text is very well written.

Potter, (1935), re-examined the microscopical features of the Hodgkin lesion; her additions of interest were a fuller description of the typical multinucleate giant-cells for which she suggested abandonment of the older eponymic titles and substitution by 'Hodgkin giant-cells'; she also gave interesting characterisations of the early stages of the disease.

Many reviews of large series of cases appeared in the latter part of this period. Relatively little of factual importance was added in respect of the typical disease. (These studies will be further alluded to later).

The Nature of Hodgkin's Disease

There is still widespread indecision on the nature of the disease. While this situation is representative of much authoritative orthodox thought, several hypotheses exist. The main theories advanced are that it is either an inflammatory lesion, or a neoplasm, or a reticulosis. In addition several less accredited ideas are, or have been, entertained. A later part of this present work is devoted to support of one of the main contentions listed, here a brief summary of the chief theories is given.

1. The Theory that Hodgkin's Disease is Inflammatory in Character.

This view survives despite a protracted series of failures to prove it exclusively tenable. Inherent in it is the interpretation that the
lesion is a granuloma, which in turn derives from clinical and pathological observations. Much of the evidence adduced in favour of this hypothesis is actually negative and consists of observations against accepting the lesion as tumour. The view is supported by many capable authorities and an enormous, if diffuse, literature. It is not practicable to construct a chronological narrative of the development of the inflammatory thesis because, till comparatively recently, systematic additions to, and analyses of, the evidence have been wanting. The idea has persisted almost since the discovery of the disease; the best considered appraisals of it are found in reviews and books devoted to the subject. In the former class the works of Symmers, (1924), and Krumbhaar, (1934), are highly critical; and in the latter the texts of Chevallier & Bernard, (1929), and Jackson & Parker, (1947), are among the best. A further valuable contribution is made by Ewing, (1929), though not as a special monograph. The arguments brought forward are conveniently listed under the headings of clinical findings, morbid anatomy, and bacteriological results.

In the first category it is pointed out that many cases are encountered in young adults and children, eras in human life in which neoplasms are uncommon. The greater incidence in males is in keeping with the distribution usually observed in infective disease. Fever, a common accompaniment of infective illness is frequent, notably in cases with extensive lesions and in the later stages. This has long been known; Murchison, (1870), first described the peculiar protracted relapsing fever characteristic of the disease. Severe sweating may be observed, and tachycardia disproportionate to the temperature rise is common.
The secondary anaemia and wasting which develop as the disease progresses are also seen in chronic infective conditions such as tuberculosis. In common with many chronic infections there may be spontaneous remissions and exacerbations throughout the relatively protracted course of the illness. The invariably fatal outcome is not inconsistent with inflammation since chronic infections like leprosy and torulosis are very lethal. Hodgkin's disease may present in an acute form, where lymph node enlargement is generalised from the start and toxic manifestations are conspicuous. In these the similarity to virulent infection is reasonably parallel.

The morbid anatomy of Hodgkin's disease unquestionably presents abnormalities which preclude a straightforward acceptance of it as tumour. The restriction to lymphoid tissue is remarkable. There is a lack of the indiscriminate metastases which characterises sarcoma, the growth is only feebly aggressive, and its rate of proliferation is relatively slow. Effusions into the serous sacs are very commonly seen but rarely are they haemorrhagic like those in neoplastic disease. At microscopical level it is held that despite the pleiomorphism of the component cells there is a degree of orderliness foreign to tumour and that the anaplasia is limited. It is also pointed out that the component cells are for the most part identical to those which partake in inflammatory reactions. In the early stages hypervascularity is often discernible but regresses as fibrosis develops, which is parallel to an inflammation with residual scarring.

The disease has been submitted to bacteriological study in a most exhaustive manner. At the outset it is pertinent to emphasize that
lymph nodes lie in the lymph stream and among their functions is the filtration of this fluid which is probably seldom sterile. Bloomfield, (1915), drawing attention to this fact, examined lymph nodes from presumed normal cases, instances of Hodgkin's disease, and miscellaneous diseases. His findings fully justified his scepticism regarding the significance of positive cultures from nodes, and his further investigations of the microbes recovered failed to identify any pathogens. He considered that morbid nodes were more likely to afford resting places for fortuitous contaminants, and this view was reinforced by Symmers, (1924).

Since Delbet, (1893), first isolated 'Bacillus Hodgkini' from a single case, and claimed to have reproduced the disease in a dog, several groups of organisms have enjoyed short-lived reputations as the casual agent. Proescher & White, (1907), found a spirochaete in four out of five cases of the disease. This microbe, Spirochaeta lymphatica, was soon abandoned however, as successive attempts at confirmation failed. From 1913 to 1916 diphtheroids were reported by Negri & Mieremit, (1913), Bunting & Yates, (1914), and Torrey, (1916). Bunting and Yates produced lesions in Macacus rhesus monkeys with their Corynebacterium Hodgkini but the similarity was not close to the human one. Cunningham, (1917), and Twort, (1930), reinvestigated the role of diphtheroids very critically and may be said to have disproved the theory beyond doubt.

Another microbe which engaged much attention for several years recently is Brucella. This work was first done in North Carolina, U.S.A. by Parsons, Poston, & Wise, (1939). Initially confirmation appeared to be forthcoming, but soon it was appreciated that this was
local only. Jones, (1946), pointed out that the localisation of the phenomenon was significant because brucellosis was endemic in the same areas. The view is now fairly generally discredited.

In 1922 Kofoid, Boyers, & Swezy, advanced the view that some of the morbid mononuclear reticulum cells in the lesion of Hodgkin's disease were in fact examples of *Endamoeba dysenteriae*. In the recognition of this identity morphology was the sole criterion adopted and the original communication related to two cases. Confirmation has not been forthcoming, and the interpretation has very little in its favour.

Simple fungi, dimorphic micro-organisms, and yeasts, have also been isolated from Hodgkin's disease lesions. In some cases full investigation including tests of pathogenuity have revealed them to be saprophytic contaminants e.g. the series reported in the Rose Research on Lymphadenoma, (1932). On the other hand examples of torulosis lesions comparable to those of Hodgkin's disease have been found, most cases are reported from U.S.A. Professor Symmers (1953) related to me a case of presumed fungal infection which produced an histological picture of the true disease so perfectly that all the members of a panel of morbid anatomists concurred in its recognition. The mycelium was cultured on one occasion and the patient has survived for longer than compatible with Hodgkin's disease.

Fitchett & Weidmann, (1934), discuss the role of torula critically and while nothing definite can be proved it is clear that torula can sometimes mimic the lesion very closely in lymph nodes. The relationship of tuberculosis with Hodgkin's disease has been examined fairly consistently over the greater part of this period. The amount of work devoted
devoted to this problem alone is prodigious. The association of tuberculous infection with Hodgkin's disease was first emphasized by Sternberg in 1898, and he initially regarded the latter as a peculiar form of tuberculosis. This view he later abandoned but the interim period being long - nearly forty years - it prompted very numerous attempts to incriminate Mycobacterium tuberculosis as the agent. It must now be conceded that tuberculosis is not an aetiological factor. For a short while the work of L'Esperance, (1929), (1931), revived the theory by suggesting that avian strains of M. tuberculosis were responsible for the lesion. General failure to confirm her findings and the reminder by van Rooyen, (1937), that avian infection though rare in man produces the usual form of tuberculous lesion have abrogated the theory. That there is an association of the two diseases is unquestionable and the many ingenious hypotheses to explain it are discussed by Wallhauser, (1933). In general terms the most reasonable conclusions are that firstly tuberculosis is a common infection, and secondly that a patient with Hodgkin's disease is more susceptible to it. This proposition may with propriety be related to the progressive reduction of physiologically normal lymphoid tissue and lymphocytes which is a definite result of Hodgkin's disease.

Up to the present there has not been any convincing evidence that a virus is responsible. Gordon, in 1932, discovered that if lymph nodes affected with the disease were suitably emulsified and injected intracerebrally into rabbits the animals developed an encephalitis. Serial transmission of the postulated virus proved impossible and subsequent
work, Friedemann, (1934), Edwards, (1938), has indicated that the con-
cephalitogenic agent is not a virus. It is noteworthy that up till his
death Gordon and some of his associated were reluctant to abandon the
virus hypothesis.

In 1950 Debré and his associates described an interesting condition
under the attractive title of 'La Maladie des Griffes de Chat'. This
newly-described disease is characterised by the development of a primary
lesion in the skin; the result of trauma and presumably infection,
inflicted by the cat. (It is also recorded that in some cases plant
thorns were the vector). The regional lymph nodes enlarge and in the
earlier stages may present a microscopical picture closely resembling
Hodgkin's disease. Further work by Kollaret, Reilly, Bastin & Tournier,
(1950), has demonstrated sensitization phenomena which appear specific
but its viral nature is unproved. The disease runs a brief course
terminating with spontaneous resolution, on occasion abscess is
developed before this; it is essentially benign.

Hodgkin's Disease Regarded as Neoplastic in Character

This view is undoubtedly gaining ground against all the others.
Progress has been gradual, every gain has been small and has had to be
consolidated. The hypothesis is as old as Hodgkin's discovery but the
means to support it have been slow in developing. At the outset it is
admitted that there is not any single great decisive fact which proves
the case to the satisfaction of all, rather are there many fragments of
evidence which collectively can resolve the lesion as neoplasm. Even
then the decision resembles a mechanism of conversion, Sargent (1951), it
/requires
requires intuitive belief. After its adoption a retrospect at the evidences and their corollaries confirms the decision and rationalises the difficulties.

The first supported contentions for tumour status for the disease were made in the opening years of the period and now at the close of it endorsement is accorded by many pathologists including some of the leading oncologists of the present day.

The facts collected in favour of the theory derive from extensive studies covering a wide field. They include the clinical and morbid anatomical findings in the common form of the disease and additional facts bearing on the problem which are latent in unusual forms of the disease or in diseases having affinities with it.

Many clinical aspects of ordinary Hodgkin's disease are compatible with the tumour hypothesis. The disease is commoner in males in the proportion of approximately 2:1., in children the ratio is even more pronounced in favour of males, e.g. Smith, (1934), submits the ratio 4:1. This disproportion conforms to analogous figures in neoplastic disease. Age is a more capricious factor. In most series recorded the incidence in children and youths is notable and most figures stress the vulnerability of these eras. This impression is not supported by all observers. Gall & Mallory (1942), reporting a large series gave the average age at initial biopsy as 36 years and Willis (1949), decided that the disease is commonest in the fourth and fifth decades of life. Geographically the disease is widespread. It appears commoner in highly civilised communities but this is probably due to the fact that its certain diagnosis is a sophisticated attainment. Typical
cases have been reported from Nigeria. Garven (1952), however, encountered very few cases during a prolonged stay in North China. Epidemiologically it exhibits the apparently fortuitous incidence of malignant disease, familial incidence is extremely rare. There is not any doubt that the disease is invariably fatal though intercurrent illnesses or complications may precipitate death. Untreated the disease runs a progressive course which seldom exceeds three years, individual cases may be very brief or unexpectedly protracted. It is attended by localised tumour, the development of weakness and secondary anaemia and in the late stages by cachexia. All remedies devised so far appear merely palliative in effect, its response to therapy being closely parallel to that of inoperable malignant disease.

Features in the morbid anatomy which favour the tumour theory are many. Those which have been made the subject of personal study will be amplified later. Here a synopsis of better authenticated facts is offered, together with observations, and arguments to meet some of the objections.

The first deliberate contention that Hodgkin's disease is a tumour was made by Gibbons in 1906. Study of this text reveals that although the writer failed to substantiate all of his thesis, he was among the first to recognise the origins of the new tissue and its affinities to lymphosarcoma. He considered it arose by proliferation of cells of the germinal centres, the endothelium lining the sinuses, the fibrous tissue cells of the capsule, blood vessels and node framework. (Ginsberg (1934), quotes Gibbons as employing 'reticulum cells', this is erroneous, the term fibrous tissue was employed). Three of Gibbons' cases were
Hodgkin's disease with malignant features, in which unusual aggressiveness was displayed by the growth. He observed rupture of node capsules and new concentric formation opposite the breaches. He also drew attention to localised outgrowths of the new tissue from the periphery of nodes which tissue was confined by capsular tissue drawn out with it. Gibbons regarded the extra-nodal deposits as metastases, but did not offer convincing evidence of their transit. On the other hand he emphasized their lack of encapsulation and distinctly infiltrative tendencies, especially in the lung parenchyma distant from the bronchial tree. Read in the light of present day knowledge, Gibbons was far ahead of his time. He also deserves credit for emphasizing the feature of capsular thickening which Reed (1902), failed to appreciate. At this period lymphosarcoma was probably a larger category than it is now. (It is interesting to note that as early as 1893 Kundrat had attempted to separate Hodgkin's disease from lymphosarcoma, but his differentiation was very poor.

Coley (1908), supported the tumour hypothesis emphatically but his reasons were confined to generalisations. These included apt comparisons of the clinical features of the disease with lymphosarcoma, and insistence on the difficulty of separating some cases from examples of sarcoma of lymph nodes.

In 1911 an important side-issue came into being, namely Hodgkin's sarcoma. This influenced the development of the tumour hypothesis; more account of it will be given later.

Oliver, (1913), produced a well-reasoned discourse on the nature of the disease and introduced the unitarian idea of the origin of endo-
thelioma of lymph nodes, lymphosarcoma and Hodgkin's disease. Strongly influenced by the contemporary work of Ewing on the foremost subject, she interpreted all three as different expressions of one neoplastic process.

Eight years later Mueller, (1921), reopened the question. Two special cases from a series were described. These illustrated the principle of mutation of microscopical picture in lymphoid tissue tumour and furthered the ideas of Coley. In his conclusion Mueller wrote 'Malignancy as it expresses itself in the destructive invasion of the neighbouring tissue and in metastases, is in these chronic diseases of the lymphatic system apparently not connected with a particular anatomical structure'. This observation reflected a mature appreciation of the peculiarity of lymphoid tissue tumour which is insufficiently advertised.

In 1931 Medlar gave a new and unexpected interpretation of the tumour hypothesis. This publication has bewildered most readers and some critics have even suggested that the author, who is no mean authority, meant it to be provocative. Briefly his conclusions were that Hodgkin's disease was a tumour arising from megakaryocytes and that the complex of cells characteristic of the growth represented a developmental cycle of the type-cell. Acceptance of the interpretation necessitated a complete reorientation of ideas about the disease, and while a new approach to an old problem is a salutory procedure this one has a speculative ingenuity verging on the incredible. By Medlar's theory primary malignant tumour arises in the megakaryocytes - a megakaryoblastoma - and metastasises to the lymph nodes and other tissues outside the bone marrow. A fundamental difficulty to most pathologists
is the morphological dissimilarity between the megakaryocytes and the Hodgkin's giant-cell. In addition the almost exclusive location of metastases in lymph nodes and gradual successive involvement of the node groups are very difficult to explain.

The material from which this study was derived deserves criticism. A considerable background of experience of Hodgkin's disease, human tuberculous lymphadenitis and of experimental work were interpreted on the basis of the findings in a single biopsy specimen of a skin nodule from a case of chronic myeloid leukaemia. This latter specimen was certainly studied intensively, but it is questionable whether this was adequate for the inferences drawn. Medlar was prepared to accept lesions without fibrosis or eosinophile infiltration as examples of the disease. This dilution of criteria is open to question; Hodgkin's disease without fibrosis is very dangerous ground. On the other hand in Medlar's favour is the fact that it is a practical impossibility to examine all the bone marrow in a human body, a postulated latent primary is therefore impossible to refute.

This tumour theory has not attracted confirmatory work.

In this year also Warthin produced an important paper which enlarged and consolidated the concept of lymphoid tissue sarcoma. The material for this study comprised 506 cases represented chiefly by Hodgkin's disease including atypical forms of it, together with examples of lymphosarcoma, leukaemia, mycosis fungoides and a small group of unclassified cases. The neoplastic status of Hodgkin's disease was expounded with emphasis, and Warthin insisted that the allied disorders named above were closely related to it. (The further contents of this
Fraser & Mekie, (1933), in a curiously written paper — it reads like a sermon — also accorded the rank of tumour to the disease, but new matter was not added to the argument.

A further study which included an excellent critical analysis of previous work was submitted by Ginsberg, (1934). The writer collated a variety of evidences of the biological similarities which could be demonstrated between Hodgkin's disease and lymphosarcoma. He also emphasized the importance of crediting reports of confused diagnosis, as these served as evidence of interchangeability of the lesions. His own work was an apt comparison of the remarkable similarities presented in the course, spread and morbid anatomy of two cases. In each instance the patient was a middle aged woman. In one patient the lesion was Hodgkin's disease throughout, in the other a large-celled lymphosarcoma.

In the last decade up till 1950, several important works have appeared which directly or indirectly support the tumour theory. These include the publications Warren & Picena, (1941), Gall & Mallory, (1942), Herbert, Miller & Erf, (1945), and Custer & Bernard, (1948). The first paper was essentially a description of the variants of sarcoma arising in lymphoid tissue, emphasizing their extremely close kinship. This schema did not include Hodgkin's disease; but Gall and Mallory integrated the latter into the generic group lymphoid tissue sarcoma. Their authors maintained at the outset that constancy of the individual members of the group was a characteristic feature, but at the same time they recorded that in 84 of their cases with serial biopsies, the original structure was maintained in 77 per cent of cases and showed de-

/differentiation
differentiation in 23 per cent. They wrote 'it is believed reasonable to expect this degree of de-differentiation in any group of cytoma followed over an extended period of time. In this particular instance the occasional transition from one form to another is a feature lending credence to the belief that these tumours are essentially of tumour origin'. Herbert, Hiller, and Erf, reported six cases in which the transition phenomenon was unusually well substantiated and demonstrated. The final work quoted provided additional evidence of the same character on a very large scale. The material studied covered 700 cases of lymph node tumour. Their results revealed that 39 per cent of the cases exhibited mutation of histological picture in the period studied.

The following diagram summarized the findings.

![Diagram of tumour types]

Experimental investigation of the disease will receive full attention in part IV. Very little has been found out however which indicates the nature of the lesion. Laser, (1951), investigated the metabolism of Hodgkin tissue and reported that it showed a type of metabolism characteristic of but not specific for tumour growth.
Hodgkin's Disease Regarded as a Reticulosis

Piney, (1925), was the first to suggest that Hodgkin's disease was an example of reticulosis, (reticulo-endotheliosis). In 1932 Pullinger developed this theory and modified the name to fibro-myeloid reticulosis in order to emphasize the predominant differentiation tendencies in the growth. Induration by fibrosis is extremely characteristic of the lesion but the myeloid affinities are less emphatically inherent. Pullinger observed the development of eosinophile granules in lymphocytes in the lesion and was able to identify progressive development of this phenomenon till cells morphologically the same as eosinophile leukocytes were produced. This she interpreted as a local myeloid metaplasia and supported the argument by assigning to the neutrophile leucocytes of the lesion a possibly similar local origin. The former metamorphosis has precedents; the formation of eosinophile leucocytes from lymphocytoid cells was demonstrated by Opie, (1904a, and 1904b), and by Dominici, (1909). It is curious that confirmatory reports of Pullinger's work on the eosinophiles of the Hodgkin's disease lesion have not been offered. Personal studies on this problem will be given later. Here it suffices to state that I have been able to confirm this transformation, but only in relatively very few samples of the disease. Further, topical eosinophilia is not invariably demonstrable in Hodgkin's disease. The eosinophiles may be so few that their presence can hardly be significant. Ross, (1933), supported Pullinger's thesis but relegated Hodgkin's disease to the 'unrestricted differentiation' category in her scheme. Robb-Smith, (1938), adopted Pullinger's name for Hodgkin's disease. He included the malady in his reticulosis classifi-
cation but, significantly, he admitted that it had features anomalous among the other reticuloses. Since the lesion usually presents first in the pulp of the lymph node it was classed as a medullary reticulosis. Among recent writers Hadfield, (1949), has supported the reticulosis theory.

In this country the reticulosis concept was quietly accepted by most pathologists and clinicians, and in many continental countries similar or more enthusiastic approbation has been given to it. In contrast the idea has never been popular in the U.S.A. Lately there has been a partial retreat from reticulosis. This is because its initial exploitation was too rapid and uncritical, and because of tendencies to fit the diseases to the theory instead of the reverse.

Other Comments on the Nature of the Disease

At the outset it was stated that a wide body of opinion existed which was non-committal on the nosological category of the malady. Even though ultimate indecision is expressed it is usually possible to detect bias in favour of one particular theory in the writings of contributors in this group. It is my impression that most of the more recent authoritative reviews give timorous support in favour of inflammation. This trend is still apparent but prior to this period tumour status was favoured more definitely, particularly in continental works.

In conclusion mention is made of several very poorly accredited theories regarding this disease. They have very little claim because most are merely guesses. It has been suggested that it is an inflammatory manifestation dependent on allergy, that it is a metabolic
disease, that any chronic inflaming agent can cause it, and that it is a unique morbid process. The evidence is simply the written word.

The Classification of Hodgkin's Disease

In all fields of study attempts to catalogue the material have a strong appeal. The purpose is to group the objects of study within a defined field so as to produce a precise and rational separation which has true value. Ideally classifications should exhibit mutually exclusive classes which are each defined by unique characteristics.

In biology it is difficult for these postulates to be met because the data are usually very complex and often incomplete. Thus most classifications in pathology are necessarily provisional, but any which may be advocated or adopted should have justification beyond being merely elegant refinement.

There is definite vindication for attempts to classify the lesions of Hodgkin's disease, even if it is adequate to regard it as an exclusive category in itself for clinical purposes.

The main support for this endeavour is that any procedure which improves the diagnosis of Hodgkin's disease is worth while. The disease is a very serious one and it is a fact that its microscopical diagnosis is not particularly well performed. Its comparative rarity determines that perfectly typical examples are not commonly seen. A classification enforces closer inspection of specimens and a better appreciation of the latitude of appearance which the lesion may present. The disease is thus better defined and other diseases are less likely to be confused with it. Failure to recognise Hodgkin's disease in its less familiar forms is quite common, yet this is a fault which is
usually corrected later; on the other hand mis-identifying it is also not rare, and is probably a more serious accident because the modern treatments of irradiation and cytotoxic drug therapy are in themselves potentially nocent.

Less obvious advantages may also derive from a classification. In time experience may relate certain variants to prognosis and likely reaction to treatment. The stability, mutation, or progression of a lesion may all be more accurately assessed when variants are recognised; and the mind becomes better conditioned to exploit the chance anomaly which is a potential key. Unexpected and interesting side-issues also emerge from this type of work, e.g. In my own studies I have encountered examples of cancer mimicking Hodgkin's disease almost to perfection, a circumstance of some importance.

Finally in attempts at classification it is common for 'scrap-heaps' to develop. These indeterminate groups though often immediately embarrassing can form the basis for new knowledge. The doubtful lymph nodes are among the most terrible treasures the morbid anatomist collects in his sojourn. Their ability to stultify the worthiest efforts, whilst humiliating, is a salutory reminder that the subject is by no means worked-out.

A really comprehensive classification of the disease is a prodigious task. It has been attempted, and setting aside the very reasonable objection that it is easy to criticise, it must be conceded that the results have not been very helpful. Ziegler, (1911), attempted it and described nine varieties. The subdivisions however are unsatisfactorily vague, being partly anatomical, partly clinical and partly microscopical.
microscopical. In several the evidence of identity is so meagre that they are almost hypothetical, e.g., the splenic and osteitic forms.

On the restricted basis of microscopical structure several tentative classifications have been evolved. Besides formal classifications, many writers have offered tentative schemata. Andrews, (1902), recognised hard, soft and intermediate varieties; Gibbons, (1906), referred to malignant Hodgkin's disease as well as the ordinary form; and Welch, (1910), first described Hodgkin's sarcoma. (This latter will be fully discussed later, it is noted here that the term is often loosely used).

Warthin, (1931), recognised typical and atypical Hodgkin's disease, and defined Hodgkin's sarcoma. Atypical Hodgkin's disease he reported to be characterised by relative or absolute lack of Dorothy Reed cells, and he added that subsequent study of these cases often revealed development of the typical picture. Kettle, (cited by Pullinger, 1932), was wont to recognise typical Hodgkin's disease and place all examples falling short of requirement for this category in 'Hodgkin Group'. Inclusion was restricted to lesions exhibiting multinucleate giant-cells and uninuclear reticulum cells. Pullinger herself described a soft cellular variety in which Hodgkin giant-cells were sparse, the reticulum cell proliferation marked, and eosinophiles few or lacking. She noted, moreover, that areas of this character were often seen in otherwise typical lesions. Jackson, (1937, 1939), submitted a simple classification which has been widely and rather uncritically adopted. He contended that the disease might present in three forms viz. Hodgkin's paragranuloma, Hodgkin's granuloma and Hodgkin's sarcoma. In the
initial description the first form was described as affecting single or multiple nodes. The predominant cells were small lymphocytes, and few or rarely many 'Sternberg-Reed' cells; eosinophiles, fibrosis, and necrosis being absent. In later publications with Parker, (1944, 1949), the descriptions differ a little, chiefly by conceding that fibrosis occurs. This microscopical entity has since been very much better described by Harrison, (1950). Hodgkin's granuloma corresponds to ordinary Hodgkin's disease. Hodgkin's sarcoma is poorly described, these authors considered it highly malignant.

Bersack, (1943), introduced another histological classification. He found he was able to segregate three varieties readily viz. Hodgkin's lympho-reticuloma, Hodgkin's granuloma, and Hodgkin's lymphoma. The first, from the description and photographs, appears to correspond with Pullinger's soft cellular variety or Gall and Mallory's stem-cell lymphoma. The 'granuloma' is ordinary Hodgkin's disease, and the last variety is like Jackson's paragranuloma. The discussion in this paper is disappointing, one has the impression it has been drastically edited. A year later Bersack advocated subdivision of the reticuloma and lymphoma groups into typical and atypical. The matter of this communication is interesting but is highly speculative.

It is common experience that varieties of Hodgkin's disease are recognised which never reach the dignity of print. These include many slightly atypical examples. The experienced, having encountered them before, can allay the tyro's concern with assurance.

There is one peculiar group in this category where the diagnosis is arrived at by a kind of reductio ad absurdum procedure - 'It must be /Hodgkin,
Hodgkin, nothing else can account for this appearance'.

**Note on Hodgkin's Sarcoma**

The characteristics of ordinary Hodgkin's disease are widely agreed and well known. In contrast there is much less understanding and agreement concerning 'Hodgkin's sarcoma'. The title is not in wide use and receives doubtful acknowledgement in most standard textbooks of pathology. It is reported rarely and the descriptions of it vary considerably.

The legend has been used in different senses and often loosely. Cases of Hodgkin's disease may present highly cellular and active lesions which prompt the description 'frankly sarcomatous'; again examples are encountered where besides an otherwise typical picture unequivocal malignant characteristics are prominent. These include cases where there is active invasion and destruction of node capsules, entrance into blood vessels, and conspicuous deposits outwith the normal sites of lymphoid tissue. From the general behaviour of such examples the term Hodgkin's sarcoma has reasonable justification.

In a more restricted meaning the title is applied to a lesion which may be regarded as a further stage of, or an unusual sequel to, ordinary Hodgkin's disease. Even in this usage niceties of distinction exist and these merit further attention. Yamasaki, (1904), is frequently quoted as being the first to employ the term. This is correct but his characterisation of the lesion was unsatisfactory, the title itself being the chief claim to priority. Welch, (1910), is properly credited as being the first to give a satisfactory report of the phenomenon.

Ewing, (1929), gave some account of this original case and afforded
full recognition to the entity. A protagonist of the view that ordinary Hodgkin's disease is granulomatous in nature he put forward the ingenious suggestion that it bore the same relation to Hodgkin's disease as squamous cell carcinoma may to a lupus scar. Ewing considered that Hodgkin's sarcoma arose most frequently in the mediastinal form (Ziegler) of the disease. The structure of the tumour varied from a close counterpart of the ordinary disease to a growth composed almost exclusively of large round cells with faintly staining granular cytoplasm, and moderately large chromatic vesicular nuclei. Large round giant-cells with multiple or multilobed nuclei might predominate in some cases. He allowed that the histological signs of malignancy were usually not pronounced, the perforation of node capsules being slow. Jackson & Parker, (1947), followed Ewing in their interpretation of the lesion. They stated that microscopically the tumour was composed of cells two to three times the size of a normal lymphocyte. The nuclei were round or ovoid and generally possessed prominent nucleoli. Multinucleate giant-cells of Sternberg-Reed type were always present. They added that besides these, scattered lymphocytes, scanty granulocytes, plasma cells and reticulum cells were also seen. The reticulin mesh was generally increased.

In a second group are those writers who, supporting the neoplastic theory of Hodgkin's disease, employ the sarcoma term in a descriptive usage for certain morphological pictures.

Warthin, (1931), considered that the sarcomatous transformation of the disease proceeded in one of two directions, to a lymphosarcoma or to a large-celled form with abundant reticulum and numerous giant-cells.
('Reticulo-endothelioblastoma'). In this latter form the lymphoid cells became reduced in number, the eosinophiles disappeared and the majority of the cells came to be large polymorphic cells with abundant cytoplasm and many large hyperchromatic nuclei. He added that numerous cells of the 'myeloid type' occurred; though exactly what was meant was not further explained. Both lesions were depicted by single photographs. Callender, (1934), characterised Hodgkin's sarcoma as identical with reticulum cell sarcoma in which pleomorphism was prominent beyond the degree exhibited by typical tumours of this cell.

The above synopses represent most of the descriptive work on the tumour.

The transformation to frankly neoplastic growth is the most significant feature. There is reason to believe that the change is commoner than generally supposed and partial transition is definitely frequent.
THE GENERAL MORBID ANATOMY OF HODGKIN'S DISEASE

There are many good analyses of the general morbid anatomy of Hodgkin's disease. Some are individual studies of small series in which the accounts are detailed, others represent the collated results of exhaustive abstracts. Representing both groups papers by the following are noteworthy: Symmons 1903; Ziegler 1911; Cunningham 1915; Symmers 1917, & 1924; Wallhauser 1933; Smith 1934; Goldman 1940; Jackson and Parker 1947; and Hoster et al. 1948.

In the following general account these works and several others will be drawn on, and supplemented by the findings in a series of 49 cases encountered at necropsy in Glasgow Royal Infirmary in the period 1900-1951.

Frequency and distribution of Hodgkin's disease

The disease is distinctly rare. In Sweden Uddstromer (1934) recorded an incidence of 0.54 per 100,000 living persons. In the U.S.A. Isaacs (1944) quoted a frequency of 1.3 per 100,000 for 1921, and 2.1 per 100,000 for 1934-6. Robb-Smith gave the figure for Great Britain (1947) as 2.3 per 100,000 living persons.

Figures from hospital mortality reports give much higher values since the population is a selected one, and this despite the fact that patients with Hodgkin's disease often die at home.
Geographically the disease has been recorded in most countries. It will be appreciated that its certain diagnosis is a sophisticated attainment, with the result that its incidence is difficult to assess in backward countries. Hoster et al (1948) record that it has been reported in Argentina, Brazil, South Africa, Australia, New Zealand and Japan. To this list may be added Nigeria, Smith (1935).

Sex incidence.

The disease is definitely commoner in males, this discrepancy being greater in children.

Age.

Figures for the age incidence shew considerable variation. Wallhauser gave the maximum in the two decades between 18 and 38 years, and...
this is probably the most general experience. On the other hand, studies of smaller series, Willis (1948), and the G.R.I. series reflect a higher incidence at a later age. In the latter the maximum fell in the fifth decade.

**Duration.**

This is difficult to determine since the onset is usually uncertain. The figures for the G.R.I. series gave an average of 2 years 2 months from diagnosis. Some authorities give a period of 3 years as average but most are agreed that individual variation is to be expected, cases as brief as seven weeks have been recorded, others as long as 20 years have been known.

In one of the G.R.I. series (P.M. 12966) the history stated that a swelling had been present in the right axilla for 20 years and that diagnosis was made from biopsy of this tumour.

**Other features.**

Congenital cases have been encountered. A well substantiated instance is that reported by Priesel and Winkelbauer (1926). Rarely it has been met with in siblings but these reports are isolated.
Tables.


Total Number of Cases. 49.

<table>
<thead>
<tr>
<th>Sex</th>
<th>Males</th>
<th>Females</th>
<th>Ratio</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>35</td>
<td>14</td>
<td>2.5:1</td>
</tr>
</tbody>
</table>

<table>
<thead>
<tr>
<th>Age</th>
<th>Decades</th>
<th>Cases</th>
<th>Sex of Cases</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>1</td>
<td>0</td>
<td>All m.</td>
</tr>
<tr>
<td></td>
<td>2</td>
<td>7</td>
<td>All m.</td>
</tr>
<tr>
<td></td>
<td>3</td>
<td>6</td>
<td>5 m.</td>
</tr>
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<td></td>
<td>4</td>
<td>8</td>
<td>10 m.</td>
</tr>
<tr>
<td></td>
<td>5</td>
<td>14</td>
<td>4 m.</td>
</tr>
<tr>
<td></td>
<td>6</td>
<td>9</td>
<td>All m. f.</td>
</tr>
<tr>
<td></td>
<td>7</td>
<td>4</td>
<td>3 f.</td>
</tr>
<tr>
<td></td>
<td>8</td>
<td>1</td>
<td>4 f.</td>
</tr>
</tbody>
</table>

Note - Relatively few children came to necropsy.

Duration of the disease from presumed onset clinically till death:

2 years 2 months (25.8 months).

(From data of 40 cases where records were adequate).
DISTRIBUTION OF LESIONS

1. Reticulo-endothelial System

(a) The lymph nodes.

It is axiomatic that lymph nodes are affected. To establish the absence of lymphadenoid lesions would require the recovery and microscopical examination of all the nodes in the body, a task which attracts few. Reasonably well accredited cases without apparent lymph node lesions have been recorded but they are very rare, some further account will be given where apposite.

Involvement is commonly assumed to be universal. This is certainly erroneous; normal lymph nodes can be found at necropsy in most cases if sought assiduously. The most likely sites include the mesenteries, the antecubital fossae and the popliteal fossae. The clinical and morbid anatomical findings often suggest that the lymph node groups become involved successively. The old observation that the disease appears first in the neck, usually on the left side, has much support, but it does not justify the postulate that the disease actually starts in this site. Symmers (1924) emphasized that deep mediastinal or retroperitoneal nodes were often conspicuously enlarged while the peripheral ones were small. In some cases all nodes appear to enlarge simultaneously (Callender 1934) but these cases are very rare and are not typical Hodgkin's disease. Restriction of involvement is commoner than might be expected, its interpretation is very difficult. The most obvious cause, namely premature death of the patient cannot always be invoked to explain it. In one case observed personally a single lymph node focus appeared to be the only lesion. (Case G.R.I. P.M.100/54).

This
This patient was an elderly man of solitary habit who was found moribund in his lodging. Necropsy revealed a large cerebral softening. An incidental finding was a solitary hard white tumour nodule in the right side of the neck which was thought to be a carotid body tumour. The lymph nodes generally were not remarkable naked-eye, none were sectioned. Microscopy of the cervical nodule revealed typical sclerosis Hodgkin's disease in a lymph node.

Affected lymph nodes are almost invariably enlarged. The increase in size is sometimes prodigious, few conditions besides Hodgkin's disease are associated with so gross an hypertrophy. Untreated, the nodes tend to remain discrete.

Different grades of maturity of the lesion are common, and alterations of the microscopical picture are also frequent. This change may amount to frank sarcomatous metamorphosis, in a later part of this work the phenomenon is studied.

Table.

Analysis of lymph nodes found to be involved at necropsy in 49 cases of Hodgkin's disease. G.R.I. series. 1900-1951.

<table>
<thead>
<tr>
<th>Site</th>
<th>Number of Cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cervical</td>
<td>39</td>
<td>80</td>
</tr>
<tr>
<td>Axillary</td>
<td>30</td>
<td>61</td>
</tr>
<tr>
<td>Inguinal</td>
<td>24</td>
<td>38</td>
</tr>
<tr>
<td>Mediastinal</td>
<td>39</td>
<td>80</td>
</tr>
<tr>
<td>Retroperitoneal</td>
<td>36</td>
<td>73</td>
</tr>
<tr>
<td>Mesenteric</td>
<td>22</td>
<td>45</td>
</tr>
</tbody>
</table>

\( \text{Table} \)
(b) **The spleen.**

In most cases the spleen is ultimately involved. Hodgkin's proposition of the sequence has been quoted, it is the general experience. The incidence of morbid change in this organ is variously reported between 56 per cent and 78 per cent of cases, Hoster et al (1948). It is clear from study of these references that distinction between enlargement and histological evidence of deposits of the new tissue is not always made. A further difficulty is that some figures are drawn from clinical studies during life.

In the G.R.I. series spleen weights were recorded in 36 cases. The average was 638 g., with extremes of 170 g. and 3,200 g. This is close to Symmers' series which gave an average of 802 g.

In 43 of the G.R.I. series the naked eye appearances were described, all shewed visible lesions i.e. 88 per cent of the full series.

The form of the lesion varies. In most studies the classical hard-bake is the commonest. Karsner (1910) however insisted that miliary deposits were the most frequent. On section the spleen is usually firmer than normal and the cut surface is more or less studded with yellowish-grey or whitish-grey nodules which vary in size from a millet seed to a walnut. These nodules have a subtle individuality. Their outlines are not so rounded as metastatic cancer nodules but are irregularly angular. They are at once incongruous. They are sharply demarcated from the red pulp, small satellite nodules may be seen round large ones. If the slice is allowed to oxygenate a deep crimson line develops round the edge of the almost white nodules producing one of the beautiful pictures of morbid anatomy.

/Necrosis
Necrosis in splenic deposits is commoner than in any other site. The remarkable appearance of the organ has prompted the true Pathologists – to whom all life is pathology (Boycott) – to draw names for it from that vital source of inspiration and simile – the kitchen. 'Hard-bake' spleen is a term of some antiquity ranking in dignity with 'sago' spleen; yet the fancied resemblance to this confection is less apt than the 'bauenwurst' of Jenz! (In Benda's house the kitchen door may have been open so that his comparison was drawn from the porphyry of the back steps).

Table.

G.R.I. Series (1900-51). Splenic lesions in 43 cases.

<table>
<thead>
<tr>
<th>Form of lesion</th>
<th>Number of cases</th>
</tr>
</thead>
<tbody>
<tr>
<td>Miliary nodules.</td>
<td>3</td>
</tr>
<tr>
<td>Solitary or scanty large nodules.</td>
<td>4</td>
</tr>
<tr>
<td>Typical Hard-bake.</td>
<td>36</td>
</tr>
</tbody>
</table>

It has been held that the disease may occur solely in the spleen or at least arise primarily in it. Exclusive involvement has been reported by Symmers (1909), Ziegler (1911), Wade (1913), and Isaacson, Spatt and Grazzel (1947). None is really satisfactory. Ziegler's contention was not properly supported. Symmers' case and Wade's case both presented with splenomegaly during life; Hodgkin's disease was discovered to be the cause, but in neither did the patient come to necropsy. The last reference is a more satisfactory account but necropsy revealed doubtful lesions in the liver and definite ones in the bone marrow.

(c) The bone marrow.

It is difficult to assess the frequency of Hodgkin's disease lesions
in the bone marrow. Complete examination of the bones is a practical impossibility in ordinary necropsies. Steiner (1943), in support of the interpretation that the disease was a reticulo—endothelial disorder rather than a lymphoid tissue one, insisted that marrow lesions were commoner than supposed. He supported this by many high figures drawn from the literature. Figures recorded vary between 6 per cent and 23 per cent but it has been postulated that the upper limit would be nearer 100 per cent if search were adequate. It is the opinion of most observers that bone marrow lesions are relatively late in occurrence. In practical study of this problem it is pertinent that the lesion can be extremely difficult to identify with certainty microscopically. Evidence of secondary anaemia is common and at necropsy extensive gelatinous degeneration is a feature in many cases.

It is almost certainly exceptional for the bone marrow to be replaced by Hodgkin tissue to a significant degree.

(d) The liver.

This organ is fairly commonly affected. Collated statistics give a frequency between 38 per cent and 60 per cent. In the G.R.I. series the data are unfortunately incomplete. In 30 cases weights were given and yield an average of 1868 g. According to Shennan (1927) the average weight for the liver is 1600 g., thus the increment is about one-fifth. Visible deposits of the new tissue were seen in 29 cases (59 per cent), in the remaining 20 cases there was not any record.

The appearance of the deposits is similar to the splenic lesions but necrosis is rare.

In small microscopical deposits the location is almost invariably in the portal tracts. This suggests that these loci favour origin of
the new tissue, and argues against the reticulo-endothelial cells of
the liver (Kupffer cells) being the originating cells, otherwise a mid-
zonal or central initium would be equally represented.

The other systems.

The lesions of Hodgkin's disease have been reported in almost every
organ and tissue of the body. It is very probable that the deposits
develop in pre-existing lymphoid tissue in these sites, but this tissue
itself may be ectopic. Ectopic lymphoid tissue is common, and may be
developed as a compensatory formation when the normal lymphoid tissue
is diseased or ablated. There is evidence that this does not readily
occur under experimental conditions (Saunders and Florey 1940). In
Hodgkin's disease the morbid encroachment is relatively protracted and
occurs in a measured piece-meal fashion; this time factor favours the
chances of development of new lymphoid tissue.

The table appended below summarises the location of lesions
outwith the reticulo-endothelial system encountered in the G.R.I. series.
Table.

Organs or tissues found to be involved at necropsy in 49 cases of Hodgkin's disease. G.R.I. 1900-1951.

<table>
<thead>
<tr>
<th>Organ or Tissue</th>
<th>Number of cases</th>
<th>Percentage</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymph nodes</td>
<td>49</td>
<td>100</td>
</tr>
<tr>
<td>Spleen</td>
<td>43</td>
<td>88</td>
</tr>
<tr>
<td>Liver</td>
<td>29</td>
<td>59</td>
</tr>
<tr>
<td>Bones</td>
<td>19</td>
<td>39</td>
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<tr>
<td>Stomach</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Small bowel</td>
<td>2</td>
<td>4</td>
</tr>
<tr>
<td>Gall bladder and ducts</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>Peritoneum, omentum.</td>
<td>3</td>
<td>6</td>
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<tr>
<td>(Ascites)</td>
<td>4</td>
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<td>Bronchi)</td>
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<td>Lungs</td>
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<tr>
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<td>Diaphragm</td>
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<tr>
<td>Kidneys</td>
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<td>16</td>
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<td>Ureters</td>
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<tr>
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<tr>
<td>Veins</td>
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<tr>
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<tr>
<td>Thymus gland</td>
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<td>2</td>
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<tr>
<td>Adrenal gland</td>
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<tr>
<td>Dura mater</td>
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<td>2</td>
</tr>
<tr>
<td>Brain</td>
<td>1</td>
<td>2</td>
</tr>
</tbody>
</table>

Note on findings in the other systems.

**Alimentary system.** The liver excepted, this system is relatively immune. Lesions are reported in the tract in 1 to 5 per cent of cases, the majority are gastric. The deposits usually commence as sub-mucosal nodules which mature to form malignant ulcers. Microscopically these lesions are usually more cellular than nodal ones. Ascites is common.
but rarely is the effusion haemorrhagic.

**Respiratory system.** Involvement of the bronchi, lungs and pleurae is relatively common, sometimes this is obviously due to extension from mediastinal nodes. Figures from the literature vary from 15 to 33 per cent. Effusion is very common, and like the ascites in this disease, blood is rare in the fluid. Extension of the disease from the diaphragmatic pleura into the diaphragm is common, in fact this is the chief expression of infiltration of muscle observed.

**Genito-urinary system.** Renal deposits are reported in 8 to 16 per cent of cases, even the lower frequency is in marked contrast to the incidence of cancer metastases in renal tissue. Ureteric involvement is rare.

**Cardiovascular system.** The pericardium is probably the commonest site, occasionally with extension into the myocardium. Invasion of veins has been reported (Ross 1933).

**Endocrine system.** Reticulo-endothelial tissue is located in the vessels of the hypophysis cerebri and adrenal glands. However the incidence of deposits in these is low. Thymus is commonly reported in children as a locus, but rarely in adults.

**Integument.** Deposits are unquestionably rare although café-au-lait pigmentation and pruritus are common.

**Central nervous system.** Deposits are very rare in the brain, less so in the meninges. (In the G.R.I. case reported extension was from the nose).

In the preparation of the foregoing survey it became obvious that there was much variation between the standards of different pathologists.
These discrepancies inevitably modified the results. As one becomes increasingly interested in a subject one's standards improve and one becomes correspondingly critical. The satisfactory antidote to this enhanced scepticism is to perform the task oneself. This has been done and the following sections embody the work and results of this endeavour.
This classical advice is difficult to follow. There are many practical obstacles to the management of large series and the ultimate analyses, whilst ostensibly objective are dependent in some measure on subjective discrimination. Full appreciation of the morphological affinities of Hodgkin's disease would require a comprehensive series of primary lymphadenopathies, entailing a formidable amount of material and study; attainments beyond the reach of those whose research is necessarily a spare-time occupation. The present series is an attempt to implement Morgagni's advice so far as it has been possible for me to accomplish. The series is primarily a collection of biopsy preparations of Hodgkin's disease. It includes less familiar and atypical forms of the lesion, and also a few cases of various lymphoid tissue tumours in which I attempt to discern some homology with Hodgkin's disease. In addition an inevitable miscellaneous group is included; this embraces lymphoid tissue disease of indeterminate nosology, remarkable reactive changes in lymph nodes, metastatic carcinoma masquerading as Hodgkin's disease, and several unsolved problem nodes.

The collection.

Altogether a total of 222 cases were studied. The intention was to restrict each serial number to an individual patient. In most cases including those with serial biopsies or later necropsy, this was /effected.
this was effected. In three patients however serial biopsies have been
given separate serial numbers viz. Ross; 71 and 76, Topping; 75 and
81, and Finnigan; 73 and 135. The series was almost complete when
these irregularities were discovered and they have been allowed to stand
so as to avoid re-numbering so many protocols blocks and slides.

Sources of the collection

Serials 1-58 inclusive refer to a collection built up by my father,
the late Professor John Shaw Dunn. These slides were found in the
house after his death and were passed to me by my mother. The
information available for most is restricted to identity and occasional
abbreviations on the labels. Serials 1-19 are from patients at the
Western Infirmary of Glasgow in the period 1910-1913. The remainder
are derived from Birmingham, Manchester and several other hospitals.
(Several private cases and some of unknown origin are interpolated
among this hospital series).

The serials 59-210 were collected by myself. These are chiefly
from the morbid anatomy routine of Glasgow Royal Infirmary but some are
from outside.

Note on the material and methods

Many of the serials 59-222 were prepared by myself. For fixation
formol-corrosive, saturated corrosive sublimate, Heidenhain's Susa, or
Bouin's solution with reduced acetic acid (1 per cent instead of 5 per
cent) were used. The mercury fixatives were found generally superior
for satisfactory staining but the modified Bouin's fluid was very good
and the tissue cut well. Ten per cent formol-saline or formol-water
was found to be poor, especially when the tissues were fresh and warm.
Zenker's fluid without acetic acid was tried but the well-known technical difficulties it produces reduced its value. It was found better to slice nodes at right angles to their long axes rather than adopt the orthodox method; distortion was less and the whole circumference of the capsule was more readily studied.

In all the serials 1-58 at least one section stained with haematoxylin and eosin was available and usually there were several. In the latest numbers of this group some connective tissue stains and silver impregnations were available. In the majority of the remainder sections stained by haematoxylin and eosin, van Gieson's method, and impregnated with silver were prepared. In a limited number Dominici's stain, Mann's stain, dilute eosin, Mallory's blue and orange, Masson's trichrome and other special stains were used. There was not any doubt that good haematoxylin and eosin surpassed all other methods in general usefulness. Eosin is an excellent polychrome stain and often indicated phenomena which special stains confirmed later. Van Gieson's stain was the most valuable adjuvant, it is more delicate than the aniline blue methods. Silver impregnation proved of doubtful value, it rarely added to what simpler methods already shewed, and introduced a new set of artificial standards.

For this study a foolscap pro-forma was made so as to record systematic descriptions of each specimen which would be mutually comparable. The following is a facsimile, reduced and condensed.

/Available

Capsule.  Thickening, hyalinisation, concentric re-formation, infiltration.

Fibrosis.  Trabecular, Peri-vascular, diffuse fibril, fibrillar, symplasmic transformation, coralline.

Reticulum cell proliferation, epithelioid, morbid mononuclear, multinucleate giant-cells, typical, atypical.


Mitoses.  Necrosis.

Diagnosis, affinities.

This schema was evolved after a preliminary survey and study.  The scope and meaning of the various headings will be fully explained in the analyses of the results.

These original protocols being unsuitable for presentation, comprehensive synopses of each have been made (See Appendix A.).  It will be noted that throughout the work interpretation has been governed by the hypothesis that Hodgkin's disease is the central expression of tumour in lymphoid tissue.  The same pattern of scrutiny has been applied to all specimens.  For each serial a diagnosis was recorded and, where relevant, explanatory remarks and etc. were added.  The serials have been re-examined repeatedly over the last five years and the ultimate assessments embody the results of these revisions.

Histories where available or traceable are included.  The clinical features are always of value, exactly as naked-eye appearances are helpful.  Yet it is remarkable how often the history recounted little more than a painless swelling at a lymph node site.  The fate of cases which I encountered personally were traced as far as possible but it is admitted that this proved singularly difficult.  In this institution, (Glasgow Royal Infirmary), patients pass through so many departments that follow-up attempts by one individual are exasperatingly hard.  In
fact what may be termed the administrative aspect of the series proved
the most tedious and frustrating task.

The microscopical examination of each specimen was as thorough as
possible. The slides were first examined under very low power, a 5 X
hard lens, and then under a magnification of 20 X. These low power
studies proved of value in many unexpected ways. Each section was then
submitted to scrutiny with the ordinary low and high-power objectives
using a Zeiss 4 eyepiece. To examine a whole section thus, occupied
about 2 hours. This is fatiguing, and it was found that oil
illumination was far superior to electricity for these protracted
observations. The restfulness of oil illumination is remarkable, it is
probably related to the constant slight movement of the flame which
reduces fatigue. Another old-fashioned device was helpful namely the
use of centre-stops in the condenser. These can be made to produce a
quasi dark-ground effect which helps to detect fibrillae & etc.
Filters were an aid. By the use of a pale blue filter, Wratten No.
78AA, inconspicuous eosinophiles were more readily detected, and con­
versely by using a red filter, Wratten's tricolour red, the granules of
eosinophiles disappeared permitting study of the nuclei. Defects in
staining, e.g. in old preparations, could be partly remedied on this
principle. In an effort to assess activity the numbers of mitoses were
counted in standard fields. The size of the high-power field was
adjusted with the draw tube till it was 0.1 sq. mm. Five random fields
of this size were surveyed and the average taken.

Findings and analysis of the series

The task of presenting a well-balanced and critical account of the
findings in the series has proved formidable.

Inherent in microscopy is a lack of objectivity, and there are practical limits to illustration. Truthful and convincing generalisations are therefore difficult to make and present.

Hodgkin's disease is above all capricious, the elasticity of appearance is remarkable and except for the characterization of important variants, classification is somewhat futile. In the belief that descriptions of the major variants might at least help in the microscopical diagnosis of the disease this has been done. So far as possible use has been made of existing categories, since to introduce new ones from studies of an already eclectic series is absurd. In the application of the classification evolved some overlap was common and at points decisions had to be arbitrary.

It is not advocated that the analysis offered should be adopted but rather that it should serve to indicate the appearances Hodgkin's disease may present and point out criteria and pit-falls in diagnosis. It is urged that study of the protocol synopses in Appendix A. will give the reader the best appreciation of the work.
I. **TYPICAL HODGKIN'S DISEASE**

In the following serials all diagnostic criteria are met. 1. 3. 6. 7. 8. 9. 10. 11. 18. 24. 34. 42. 56. 64. 65. 66. 69. 70. 71. 74. 75. 76. 79. 81. 83. 89. 91. 92. 99. 105. 106. 113. 114. 115. 116. 120. 122. 126. 132. 138. 184. 188. 195. 201. 211. 212. 216. 217.

Total 48 examples from 46 cases; with the exception of Serial 81, all are node lesions and represent all grades of maturity.

**Features of the typical lesion**

(A) **Habit and mode of growth**

The initium of the new tissue is commonly nodular in form, and this characteristic may persist. It is usual for it to develop in the deep pulp of the node but it is occasionally cortical in position. This primary focus of the disease is generally single. As it grows it rapidly takes up the surrounding lymphoid tissue and also displaces it; in this study this has been termed the exodic growth phenomenon. The demarcation between the periphery of the expanding nodule and the surrounding lymphoid tissue is best appreciated by low-power examination of the order 5-20 diameters. In many cases high-power scrutiny can confirm that the peripheral lymphoid tissue shews only physiological reactive hyperplasia. In 22 cases this partial or incomplete involvement of the node was a feature. The surviving lymphoid tissue may contain solid follicles, follicles with pale centres, and even sinus tissue. In many otherwise totally involved nodes small aggregates of normal small lymphocytes, survival islets, were found.

In 10 cases a further phenomenon related to this expanding nodule conformation was detected namely the development of a swathe or swathes...
of new connective tissue roughly parallel to the true capsule. This has been termed concentric re-formation of the capsule.

As the lesion matures all or almost all of the original tissue of the node is replaced by the new tissue. In these established lesions a more or less segmental or folliculoid pattern is common. This was detected in 24 examples. Ultimately the picture is one of irregular islets of the new tissue set in a coarse fibrous matrix.

(B) The new tissue

This consists of the products of proliferation and fibroblastic transformation of reticulum cells, mature and immature lymphocytes, eosinophiles, neutrophiles and plasma cells. Even in the earliest stages all components are present. It can be stated categorically that the earliest recognisable stage of typical Hodgkin's disease is the presence of this characteristic new tissue, which it has already been seen may be only focal. Pictures of reticulum cell proliferation without fibrosis, lymphoid hyperplasia, or general loss of node architecture alone are not ever safely labelled early Hodgkin's disease.

1. Reticulum cell proliferation

This is invariable and produces several characteristic cells. Sometimes it is distinctly focal, especially where the gross pattern is strongly folliculoid. The pulp reticulum cells are primarily involved but the others may partake. Four main cell-types can be described.

(a) Morbid mononuclear reticulum cells.

These are always present, they are at least twice the size of normal reticulum cells, certainly in a nuclear respect. They are characterised by large plump nuclei which are usually oval or spheroidal and possess a
well developed nuclear membrane. Nucleoli are common and may stain as plasmasomes or as Karyosomes. The chromatin is generally aggregated.

(b) Hodgkin giant-cells.

These are invariably present. Mitoses are rarely seen in these cells and it is common for them to be lost in the course of preparation. Their form is remarkably protean.

(c) Epithelioid cell-types.

These can generally be found but are sometimes absent. They correspond closely to the epithelioid cells of early tuberculous lesions or to macrophages. The nuclei are characteristically smaller than those of (a). Where present these cells are often seen in closely set clusters.

(d) Fibroblastic cell types.

These are always found. They correspond to the plump mildly pleomorphic fibroblasts of active granulation tissue, but are usually larger.

2. Fibrosis.

True collagenous fibre production is invariable. There may or may not be reticulin increase. Fibrosis is expressed in several ways.

(a) Augmentation of pre-existing fibrous tissue in the node capsule and trabeculae. It is frequently gross. The mechanism is difficult to follow but it appears in some cases to derive from cells (d) above.

(b) The appearance of fibrillae.

This occurs in the new tissue pulp. The origin of these most delicate collagen fibres may be traced to morbid reticulum cells of types (a), (b) and (d), apparently forming in the cell cytoplasm.
small number of fibrillae can be traced to type (c) cells. Here they appear as condensations of contiguous cell margins. This mechanism has been termed symplasmic transformation. It is very similar to the fibrosis process in tuberculous lymphadenitis.

(c) The development of fibrils.
(The term fibrilla is used as a diminutive). These originate as fibrillae and are regarded as condensation and accretion products of them.

(d) The formation of coralline deposits of collagen.
These may take the form of tortuous cylinders which have many outlines or of sieve-like structures resembling red coral (Corallum rubrum). They may be related to antecedent fibrils but are often seen well apart from them. Their origin is somewhat mysterious, they are very rarely of vascular source.

(c) Perivascular lamellation.
This form is very characteristic and particularly affects arterioles. In transverse section the vessel is seen surrounded by concentric, but often incomplete, collagenous lamellae.

Analysis of the studies made reveals that the most constant forms of fibrosis in the lesion are fibrillae and fibrils. In general terms the more cellular the lesion the less is the fibrosis. The remaining types are commoner in slowly evolving examples of the disease. It may be added that post-infarction scarring is occasionally seen, it appears to result from granulation tissue formation round necrotic foci, the process is rarely complete.

3. **Lymphocyte series cells**

The normal lymphocytes of the node always diminish. The decrease
may be slow or rapid and is roughly proportionate to the rate of growth of the new tissue. It is extremely common to find a proportion of immature lymphocytes, including the earliest precursors. These immature lymphocytes may virtually replace the normal ones. Their immaturity is reflected by appreciable enlargement and leptochromicity of their nuclei.

4. **Eosinophilia.**

In the typical disease eosinophiles are present in sufficient numbers to arrest attention at once. In most cases they are typical binucleate blood-stream forms. The granules are fewer and coarser but this is due to histological as opposed to cytological preparation. They are very capricious in distribution, often tending to be focal. One fairly characteristic site is in the interstices of new young connective tissue. Eosinophiles with single lymphocytoid nuclei are sometimes present. In 7 cases (14.5 per cent) these cells were demonstrable and transitions could be traced with some conviction. Special methods were essential for the purpose, the best results being obtained with Dominici's stain.

5. **Neutrophiles and plasma cells.**

These cells can always be found. Most often they are scattered diffusely through the new tissue but may form small aggregates. Neutrophiles are occasionally sufficiently numerous to indicate superadded inflammation and may also be notable round necroses.

6. **Necrosis.**

In 22 cases of the series necrosis was absent in any form. In 9 cases it was limited to scattered individual cells and in the remainder /occurred
occurred in small foci. It is generally a quiet necrosis of coagulative fibrinoid or caseous type, which evokes very little reaction.

7. Mitotic activity.

The results of this survey proved disappointing, the figures recovered pretend an accuracy which is false. In 37 cases 1 or less mitotic figures presented in each standard high power field, but in several cases where multiple sections were examined great discrepancies appeared.

II. ALMOST TYPICAL HODGKIN'S DISEASE.

Due to the strict application of criteria the following serials were excluded from the category Typical Hodgkin's disease. 14. 16. 17. 96. 98. 101. 107. 110. 121. 129. 196.

Total 11 examples from 11 cases.

It is emphasized that deviation from the characteristic lesion was minimal.

The group is constituted as follows.

Cellularity above average. 6 cases (14. 16. 17. 96. 101. 196.)

Presence of rare Langhan's giant-cells. 1 case. (98).

Large morbid reticulum cells numerous. 1 case. (107).

Eosinophiles below average. 1 case. (110).

Conspicuous epithelioid cell foci. 1 case. (121).

Anomalies attributable to site. 1 case. (129). mesenteric node.
II. **ATYPICAL HODGKIN'S DISEASE.**

This is a large category. In all 75 examples were encountered in the series. The atypism varied from comparatively minor deviations from the typical lesion, to relatively major alterations. Cases representing these extremes are easily separated but analysis of the intermediate ones is difficult because of the lack of consistent objective criteria. In general terms the whole group is considered under two main headings viz., major and minor variants. These subgroups are then divided on a basis of the most obvious mutual differences.

A. **Major variants.**

1. **Soft cellular and tumour-like Hodgkin's disease.**

In this form the lesion is still recognisable as Hodgkin's disease. In some examples this recognition depends on much experience, and in others the lesion is passing to another category. A proportion of these cases shew strong morphological affinities to other lymphoid tissue tumours.

This category is divisible into two depending on whether eosinophiles are retained or not.

(i) **Eosinophiles retained in normal or slightly reduced numbers.**

Serials. 31. 32. 40. 60. 62. 63. 87. 100. 111. 112. 118. 119. 131. 193. 198. Total 15 cases.

In serials 111. 119. 193. the lesion was cellular but not tumour-like; in serials 40. 60. 63. 100. 112. 118. 131. 198. tumour-like features were more pronounced; in serials 31. 32. 62. 87. neoplastic features were extreme.

**Features.** Persistence of normal lymphoid tissue is less than in the
typical disease but the general habit of growth is the same. The
eosinoplastic growth phenomenon is less common and so is concentric re-formation
of the capsule. The node capsule does not usually show marked hyaline
thickening and it is sometimes infiltrated with the new tissue.

Fibrosis is characteristically much less in total amount and is
expressed chiefly by fibrillary and diffuse fibril formation. It is
mainly on this decrease of fibrosis that the cellular appearance depends.

The reticulum cell proliferation is usually florid and is
productive of smaller cells than in the typical disease. Morbid mono-
nuclear cells and Hodgkin giant-cells are both affected. It is not
uncommon for this cell proliferation to be focal. Eosinophiles while
persisting tend on the whole to be reduced. The lymphocytes are
commonly large and immature in type. Plasma cells and neutrophiles are
little affected but necrosis especially on an individual cell basis is
common. Mitotic activity is either normal or increased, these lesions
are rarely indolent.

The tumour-like designation is applied where an unwonted homo-
geneity is apparent in the new tissue. All grades are encountered to a
point where it is equivocal whether reticulum cell sarcoma is not an
equally correct microscopical diagnosis. It is noted that where
eosinophiles persist metamorphosis to other lymphoid tissue tumours is
not seen.

(ii) Eosinophiles reduced or lost.

Serials. 57. 67. 72. 77. 80. 97. 102. 103. 127. 128. 130. 143. 154.

In serials. 57. 72. 80. 127. 130. 143. 154. 176. 180. 181. 205.

/
the lesion was cellular with relatively unimpressive tumour-like appearance; in Serials 67. 77. 97. 102. 103. 128. 159. 161. 185. 213. 207. 218. neoplastic features were pronounced.

**Features.** These are essentially the same as in the previous heading. It is noted that the total encountered is larger and that proportionately more of the cases qualify for tumour-like affinities. While the general direction of affinity is towards reticulum cell sarcoma other variants are seen. In Serials 72. 127. 180. 207. an approach to the microscopical features of lymphosarcoma is notable; and in Serial 80 some features of Brill's disease are observed.

2. **Diffuse fibroblastic Hodgkin's disease.**

This uncommon variant was present in Serials 124. 140. 153. 168. 219. 220. Total 6 cases.

This variant is characterised by a striking increase of fibroblastic cells, the majority of the cells present being of this character. Typical morbid mononuclear and Hodgkin's giant-cells are present but may be absent in individual sections. Eosinophilia is absent or less than usual. Small lymphocytes are scanty. In some cases whorling of the new fibroblastic tissue is a feature recalling indolent fibro-sarcoma. There appears to be little tendency for the fibrillae and fibrils to augment by accretion so that these lesions often present an openwork appearance. In two of the cases epithelioid cells were conspicuous.

This variant is treacherous, in Serial 168 the diagnosis was only made after many sections had been taken. While six cases is a very limited series it appears that these cases always do badly. They react
very poorly to radiotherapy, and most of these patients were dead within six to twelve months of diagnosis. Clinically they also presented difficulties; lymphadenopathy was often slight, and weakness was usually the chief complaint.

3. **Lymphoblastic Hodgkin's disease.**

   **Serials.** 53, 85, 158, 214. Total 4 cases.

   This is not a very satisfactory category because lymphoblastic differentiation was not the sole abnormality nor is it restricted to these cases. However in these four examples the phenomenon was pronounced.

   **Features.** Undue prominence of immature lymphoid series cells is seen in all. The appearances indicate that the reticulum cell proliferation is given over particularly to production of lymphocyte precursors. Fibrosis is slight and in all except one, eosinophilia is lacking. All are considered nearer to Hodgkin's disease than any other variant of lymphoid tissue tumour, but confusion with lymphosarcoma or Brill's disease could arise. (See protocols of Serial 85, 214).

4. **Benign Hodgkin's disease.**

   The microscopical appearances described for this entity were observed in Serials 2, 54, 137. Total 3 cases.

   In only one case, Serial 54, was a history of relatively long survival recorded. While there is little doubt that this recently described lesion is real enough it is dangerous to diagnose because it is very rare. The essential features appear to be conspicuous fibrosis of the capsule and trabeculae with resultant segmentation of the node, little if any fibril-type fibrosis and an indolent slightly immature lymphocyte.
lymphocyte population in which rare morbid reticulum cells are seen. (The relevant protocols should be consulted for details).

In Serials 23, (30), 33, lesions which were interpreted as links between typical Hodgkin's disease and the above variant were encountered.

**Features.** In these cases the fibrosis is widely scattered but not so diffuse as in the typical lesion. The reticulum cell proliferation is below average and pleomorphism is reduced. Eosinophiles are low and the small lymphocytes are mature types. (Serial 30 is anomalous, it is nearer to the typical disease but is a very early lesion).

5. **Atypical Hodgkin's disease with prominent epithelioid cells.**

**Serials.** 47, 68, 117. Total 3 cases.

In these cases epithelioid cells were very conspicuous but were not the only abnormal feature.

**Features.** In this form islets of epithelioid cells occur in patchy fashion throughout the new tissue. Symplasmic transformation fibrosis is commonly seen in relation to them. Differentiation to this cell appears to occur at the expense of morbid mononuclear and Hodgkin giant-cells. Fibrosis of the usual form is slight and eosinophiles are few. Differentiation from endothelial tuberculosis can be very difficult.

6. **Miscellaneous atypical Hodgkin's disease lesions.**

The following did not correspond to any of the variants listed above.

**Serials** 44, 59, 206.

Serial 44 represents an extraordinary example of the disease. All components of the normal complex are present but the lesion is a parody of the disease.

Serial 59 is Hodgkin's disease growing in the integument. Pseudo-suppuration is present and granulation tissue complicates the picture.
Serial 206. The appearances in this case strongly favour a sinus origin of the lesion. Morbid reticulum cell production is seen in sinus sites but is not restricted to them. (Eosinophiles are also much reduced).

B. Minor atypical features in Hodgkin's disease.

1. The absence of eosinophiles in otherwise typical lesions.

This was observed in Serials 15, 30, 70, 80, 134, 162, 176, 181, 219. Total 9 cases.

In otherwise typical lesions this anomaly does not invalidate the diagnosis of Hodgkin's disease; eosinophilia is characteristic but not essential. It may sometimes be absent at the level first examined but present in further sections. In late lesions, especially after irradiation, a marked reduction of eosinophiles is common.

2. The presence of abnormal multinucleate giant-cells.

These were seen in Serials 7, 13, 21, 53, 124, 153. Total 6 cases.

In Serials 7, 13, 21, the cells were typical Langhan's type giant-cells and constituted the only aberrant feature. Evidence of tuberculosis was lacking.

In Serial 53 the lesion was tumour-like and the cells corresponded closely to foreign-body giant-cells. In Serials 124, 153, the lesions were also atypical and the cells of Langhan's type.

This phenomenon possibly reflects the ability of some reticulum cell descendents to react physiologically, there is a little evidence that antagonism exists between the Langhan's cells and Hodgkin giant-cells.
III. FALSE HODGKIN'S DISEASE.

1. Metastatic carcinoma masquerading as Hodgkin's disease.

This was encountered in Serials 95, 174, 222. Total 3 cases.

Features. This hazard in diagnosis is rare. It is most likely to cause difficulty where carcinomatous invasion of a lymph node is diffuse, the individual tumour cells lying out of contact with one another. Topical eosinophilia may accompany infiltration of this variety and brings the similarity to Hodgkin's disease closer. Diffuse fibroblastic reaction to the invading tumour also mimics the true disease. Study of further sections is the most helpful measure in establishing the true diagnosis.

In Serial 95 the diagnosis remains open. The case has been followed for almost four years without any development of a pathognomonic evolution. This latency is unusual for primary or secondary tumour but the appearance are microscopically outwith the limits of reactive change. In Serials 174, 222. the diagnosis of metastatic carcinoma is sustained.

Note. Metastatic melanoma in lymph nodes may mimic Hodgkin's sarcoma. In these cases the tumour is unpigmented and the large cells closely simulate morbid reticulum cells.

2. Pronounced physiological reaction simulating Hodgkin's disease.

In the following serials marked physiological reactive change was encountered. Serials 84, 86, 94, 148, 152, 155, 157, 165, 171, 178, 182, 183. Total 12 cases.

Features. In the majority of these cases the mimicry is not close and is often restricted to single characteristics. In four examples however, Serials 94, 148, 155, 171, real difficulty in separation from early or atypical Hodgkin's disease is present.
In these reactive hyperplastic states there is a more or less conspicuous proliferation of reticulum cells. The majority of cells resulting have smaller nuclei than those in the proliferation characterising Hodgkin's disease and giant-cells are very rare. At the same time a few typical morbid mononuclear types are not uncommon. Attending this hyperplasia there is some general loss of definition of the normal architecture but it is never complete. Fibrosis other than capsular and trabecular thickening is unusual, and this deficiency is most helpful in differentiation. Topical eosinophilia may be present but is rarely up to the normal level observed in lymphadenoma.

In Serial 94, difficulty arises because of apparent fibrosis in the pulp of the node but this interpretation is false as the connective tissue can be traced to the hilar intrusion.

In Serial 148, the size of the node was considerably above normal which is a very suspicious feature. The subsequent history of this patient is worth relating because it justifies the rejection of the original diagnosis. After being told the nature of his malady and advised to attend for X-ray therapy, he promptly entered a recruiting office and was accepted for the Army without demur. He went overseas with his unit and served three years with the Colours in the Far East. On his return to the United Kingdom he was relegated to the Reserve and took up his old work. Through the courtesy of the family physician, Dr. Boyle, who has helped me most generously, I have been able to trace the patient to date and can confirm that he is alive and well.

Serial 155 is an interesting lesion probably attributable to X-ray therapy. Endarteritic thickening of the blood-vessels in contrast to perivascular thickening is notable.

/Serial
Serial 171 is an example of reactive change in which some follicles are very large. Superficially the appearances recall Brill's disease and changes suggestive of early tuberculous inflammation are also seen.
IV. **HODGKIN'S DISEASE CO-EXISTENT WITH OR IN TRANSITION TO RETICULUM CELL SARCOMA.**

These cases were found in the following Serials and grouped thus.

1. **Recognisable typical or atypical Hodgkin's disease accompanying some variety of reticulum cell sarcoma.**
   

2. **Lesions where a process of transition between Hodgkin's disease and reticulum cell sarcoma is apparent.**
   

3. **Hodgkin's disease co-existent with Brill's disease.**
   
   Serial 113.

**Features.**

1. The form of reticulum cell sarcoma may be either predominantly a stem-cell growth, Serials 190. 191. 210., or Hodgkin's sarcoma type 2 of Warthin (reticulo-endothelioblastoma), Serial 22. The lesions may be seen together in the same node or separately in different nodes. Transition is discernible in three cases, Serials 22. 191. 210.

2. In this group the lesion is Hodgkin's sarcoma in all the cases except one, (Serial 204), where further transition to a stem-cell type of growth was recorded later.

The most important changes in this development of Hodgkin's sarcoma are a disporportionate increase in morbid reticulum cells, repression of eosinophilia, reduction of lymphocytes and increase in growth rate. The pleiomorphism of Hodgkin's disease is retained partly.

3. This example is unique, representation of the lesions is about equal.
V. RETICULUM CELL SARCOMA.

This tumour was diagnosed in the following groups.

A. Pure reticulum cell sarcoma.

Serials. 5. 25. 26. 35. 39. 43. 48. 52. 61. 73. 78. 82. 109. 133. 135. 141. 142. 156. 221. Total 19 cases.

B. Intermediate reticulum cell sarcoma.

Serials. 12. 19. 41. 49. 50. 90. 163. 194. 200. Total 9 cases.

Analyses of both groups was made with particular reference to features in which they shewed morphological affinities to Hodgkin's disease.

Features, Group A. Folliculoid or segmental habit is seen in 7 cases but is absent in the remainder. Fibrosis in the forms displayed in Hodgkin's disease is detectable in 15 cases. In all, morbid mononuclear reticulum cells can be found, though the actual number present may be very small. Giant-cells identical to or closely resembling Hodgkin giant-cells are seen in 17 cases. Eosinophiles are present in only 1.

It is considered that the morphological affinities to Hodgkin's disease are near in 8 cases, moderate in 2 and remote in the remainder. Three of the cases shew affinity with Brill's disease.

Group B. Folliculoid or segmental habit is seen in 5 cases, and Hodgkin's disease types of fibrosis occur in 6 cases. In all, morbid mononuclear reticulum cells are detected, and Hodgkin giant-cells are encountered in 7 cases. In 3 cases eosinophiles are present but never in large numbers.

It is considered that the affinities of these growths to Hodgkin's disease are near in 4 cases, moderate in 1 and remote in the rest. Two cases shew some affinity to lymphosarcoma and three to Brill's disease.

/Note.
Note. In both these groups the affinity to Hodgkin's disease is assessed either by the sum of the corresponding features or by the finding of microscopical fields which simulate the disease.
### Analysis of 19 cases of pure reticulum cell sarcoma.

#### Biopsy Series.

<table>
<thead>
<tr>
<th>Serial</th>
<th>Hodgkin fibrosis</th>
<th>Mononuclear morbid reticulum cells</th>
<th>Hodgkin giant cells</th>
<th>Eosinophils</th>
<th>Affinity to Hodgkin's disease</th>
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**P = present**  
**N = near**  
**R = remote**  

**A = absent**  
**M = moderate**
Analysis of 9 cases of intermediate reticulum cell sarcoma.

Biopsy series.

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<th>Serial</th>
<th>Hodgkin fibrosis</th>
<th>Mononuclear Hodgkin morbid reticulum cells</th>
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P = present  N = near  R = remote
A = absent    M = moderate
VI. **BRILL'S DISEASE.**

This was recorded in the following, Serials 20. 27. 28. 36. 38. 45. 46. 51. 104. 108. 145. 172. 173. 203. Total 14 cases.

It is well-known that this entity is the least stable variant of lymphoid tissue tumour. This inconstancy evidenced by incipient metaplasia was confirmed among these examples, and in several the growth was atypical but morphologically nearest to Brill's disease.

**Features.** Folliculoid habit is present in all but early loss of the characteristic perfection is apparent in several.

In 10 of the group fibrosis is notable. This is generally perifollicular but sometimes diffuse in the inter-follicular pulp. In the other 4 the phenomenon is insignificant. Although visible in 12 of the cases morbid mononuclear reticulum cells are always few; similarly Hodgkin giant-cells are hard to find and often atypical. They are seen in 8 cases. In most of these tumours the type-cell is closely similar to that of the intermediate reticulum cell sarcoma. Eosinophiles are present in only 1 case.

Affinity to Hodgkin's disease is less in evidence. Only in 4 is it near, in 1 it is moderate and in the remainder remote or absent.

In 3 of these tumours affinities to lymphosarcoma are noted, in 1 conversion is in progress.
Analysis of 14 cases of Brill's disease. Biopsy Series.

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P = present  N = near  R = remote
A = absent    M = moderate
VII. **LYMPHOSARCOMA.**

This tumour was encountered in the following forms.

1. **Small or intermediate cell lymphosarcoma.**
   Serials. 123. 144. 149. 150. 151. 169. 170. 175. 186. 192. 197. 208. Total 12 cases.

2. **Large cell lymphosarcoma.**
   Serials. 136. 147. Total 2 cases.

These tumours have least morphological affinity with Hodgkin's disease, the characteristics that are parallel are usually minimal expressions.

**Features.** Folliculoid habit is detectable in 8, and in 3 of these the tumour is probably an evolution from antecedent Brill's disease. Fibrosis in forms characteristic of Hodgkin's disease is seen in half of the group but the total amount of collagen is small. Relatively isolated morbid mononuclear reticulum cells are present in 12 cases but Hodgkin giant-cells are only found in 6. With the exception of 1 case eosinophilia is conspicuously absent. These findings are reflected in the assessments of affinity of these growths to Hodgkin's disease, 1 shews moderate affinity, 3 remote likeness and 10 none at all.
Analysis of 14 cases of lymphosarcoma. Biopsy Series.

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</table>

A = absent  N = near  R = remote  
P = present  M = moderate
VIII. LYMPH NODES IN MYELOID LEUKAEMIA.

Two examples were encountered, Serials 179 and 189. These cases are of much interest because in both a tentative diagnosis of acute or atavistic Hodgkin's disease was made from lymph nodes before blood examination established the true diagnosis. These diagnoses were invented to indicate that the impressive features were the immaturity of the cells of the complex, and the tenuous light fibrosis. It has already been recorded that this identity between myeloid leukaemia and Hodgkin's disease has convinced Symmers and Medlar. It is ironic that I should have rejected the hypothesis of these writers and then have subscribed to it unwittingly. Further, I repeated the error within six months. The lessons to be learned are that blood examination should be made before diagnosis of unusual variants of Hodgkin's disease is completed, and that speculative morbid anatomy is unsound.

IX. INDETERMINATE LYMPHOID TISSUE TUMOUR

This unsatisfactory diagnosis was necessary in the following, Serials 29, 37, 55, 58, 88, 139, 146, 164, 166, 167, 177, 187, 215. Total 13 cases.

While the characteristics of these lesions did not permit diagnosis in existing categories, they are provisionally grouped thus:

1. Nearer to Hodgkin's disease than other variants

2. Indeterminate lymphoid tissue tumour
   Serials. 55, 58, 88, 139, 166, 177. Total 6 cases.

3. Doubtful
   Serials. 37, 164. Total 2 cases.

/Analysis
Analysis beyond this is not profitable. The original protocols should be consulted for details of individual cases.
THE NECROPSY SERIES

It is difficult to obtain necropsies in cases of Hodgkin's disease. This limitation of material enforces the exploitation of any that may be obtained, and consequently the threshold of acceptance tends to be lower than in eclectic series of commoner diseases. The present collection comprises 13 cases gathered over a period of five years. Most of these necropsies were performed personally and single-handed; in two cases I am indebted to colleagues who permitted me to re-examine the cadavera after routine necropsy. All the material from every case was processed, cut, and stained by myself. This entailed much work since each case required 50-70 blocks, many of which had to be kept mutually separate. Inevitably some material has been badly chosen, incomprehensibly forgotten, lost at the last moment, or mis-identified. Similarly as experience increased more extensive and sophisticated surveys developed. These factors diminish the completeness and mutual comparability from the ideal contemplated but not too seriously. The specimens selected for microscopy varied in freshness from 2 hours to 2 days, most were under 12 hours. The standard of histology attained was reasonably good, the exceptions where post-mortem change was advanced, shewed the usual capricious relationship to the actual time involved.

The main interests in this study were the distribution, location, and extent of the lesions. In addition the stability of the lesion and several subsidiary issues have received attention.

Material.

The series consist of twelve cases collected from the routine necropsies in Glasgow Royal Infirmary, and the Eastern District Hospital, /Glasgow,
IDENTITY OF THE CASES.

<table>
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<th>No.</th>
<th>Code Letter</th>
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General principles of selection of material for microscopy.

Lymph nodes. Representative samples from all accessible groups were taken including specimens from both sides in the case of paired groups. Where nodes were very large, pieces only were taken. Those from the face and base of the skull are difficult to extract without mutilation and attempts were limited. The nodes in the antecubital, epitrochlear and popliteal regions were investigated in all except the earliest cases reported.

Organs (chiefly viscera, but including some outwith the large body cavities).
The selection was determined primarily by the results of naked eye
scrutiny. In such viscera as lung, liver, spleen and kidney it was
found valuable to cool the tissue till it was rendered appreciably rigid
and then slice it closely. In cases where naked eye lesions were absent
random samples were taken.

**Skeleton.** Thorough examination of the bone marrow is an impossibility
in ordinary necropsy practice because it produces serious mutilation
besides being exceptionally tedious. In most cases examination was per­
force limited to the skull, sternum, ribs, bodies of the vertebrae, and
one femur. The nuclei pulposi were examined in 8 cases. The method
found most useful was to dissect the structure after it had been partly
approached by the tangential saw-cut made to examine the vertebral bodies.
(Microscopy was unsatisfactory in these structures, they almost
invariably disintegrated during processing).

**Integument and fascial planes.** The entire skin was inspected and pal­
pated, any suspicious irregularity or swelling being removed en bloc.
Examination of superficial and deep fascia was restricted to the anterior
thoracic and abdominal representative. The omenta and mesenteries and
the fascia of the lower limb opened were also examined closely.

**Methods.**

**Fixation:** In most cases 10 per cent formol-corrosive was employed as a
routine. Many hard lymph nodes however were fixed in Bouin's solution
with reduced acetic acid (1 per cent), since this was found to give
better results. Bone marrow specimens were fixed in 'Susa', the decal-
cifying properties of this acid solution being generally adequate to
soften small spicules of bone included in these specimens while at the
same time giving excellent results in staining.
Dehydration and clearing were effected with the usual reagents including toluol. The tissues were embedded in paraffin and sections 5 µ thick were cut.

Stains. All sections were stained with Harris' haematoxylin and water soluble eosin. In selected cases further sections were stained by connective tissue stains principally van Gieson's stain and Mallory & Orange G. These were supplemented by silver impregnations (Gordon & Sweets 1936) and (Mitchell & Wislocki, 1944). The latter method gives beautiful results but is expensive to use. In a few cases, where indicated, Ziehl-Neelsen's stain, Lieshman's stain and Dominici's stain were employed.

RESULTS

Morphological findings are difficult to present briefly. In each case a full necropsy report of the naked-eye and microscopical findings has been prepared; these are supplemented by homunculus diagrams recording the reticulo-endothelial system findings. In addition a series of photographs relevant to each case has been added. The full protocols are in Appendix 'B'.

Here the results are confined to analyses of the subjects referred to in the introduction above.

1. Distribution of the lesions.

To facilitate the presentation of the results of this survey each necropsy has been performed and recorded on a definite plan.

Appended is a schema which covers nearly all of the morbid anatomy studied.
DISTRIBUTION OF THE LESIONS

Schema of the basis of analysis of the necropsy series.

I Reticulo-endothelial system
   A. Reticulo-endothelial organs.
      1. Lymph nodes.
      2. Spleen.
      3. Bone marrow.
   B. Reticulo-endothelial elements situated in organs.
      1. Liver.
      2. Adrenal glands.
      3. Hypophysis cerebri.
   C. Dispersed elements of the reticulo-endothelial system.
      1. Skin.
      2. Omenta.
      3. Fascial planes.
      4. (Nuclei pulposi of intervertebral discs, persistent mesenchyme).

II Respiratory system. (excluding intra-pulmonary lymph nodes).

III Alimentary system. (excluding adnate mesenteric lymph nodes).

IV Serous Membranes.
   1. Meninges.
   2. Pericardium.
   3. Pleurae.
   4. Peritoneum, other than mesenteries and omenta.

V Genito-urinary system.

VI Cardiovascular system.

VII Skeletal muscle system.

VIII Endocrine glands. /Systematic
Systematic survey.

Ia. 1. Lymph nodes.

The results are recorded in the following table.

**TABLE SHOWING INCIDENCE AND DISTRIBUTION OF LYMPH NODE LESIONS**

<table>
<thead>
<tr>
<th>Site of Lymph node</th>
<th>A</th>
<th>B</th>
<th>C</th>
<th>D</th>
<th>E</th>
<th>F</th>
<th>G</th>
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<th>I</th>
<th>J</th>
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<th>M</th>
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<tbody>
<tr>
<td><strong>NECK.</strong></td>
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<td><strong>ABDOMEN.</strong></td>
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<td>Mesenteric.</td>
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<td>Greater omentum.</td>
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<tr>
<td>Mesentery S.I.</td>
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<td>Mesentery L.I.</td>
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<td>Retro-peritoneal.</td>
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<td>Para-aortic.</td>
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<td>L. Groin.</td>
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<td>R. Groin.</td>
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/
The term lymphoid tissue tumour covers Hodgkin's disease, Hodgkin's sarcoma and reticulum cell sarcoma. In nearly all the cases non-involvement was confirmed microscopically. In the majority of cases the nodes of the antecubital fossae could not be found; search for those in the popliteal fossae met with rather better success. In all the cases except case E. at least one and occasionally many persisting non-involved nodes were found. These nodes were generally atrophic, fat atrophy and lymphocyte depletion being very common. A number have been photographed.

I.A.2. The Spleen

In ten cases lesions were identified naked-eye and confirmed by microscopy. All the deposits were in the form of nodules of which the smallest were consistently sited in the red pulp. In the remaining three cases naked-eye scrutiny was negative and microscopy confirmed the absence of the disease. Necrosis and haemosiderin deposit were frequently notable in the foci in this locus.

I.A.3. The bone marrow.

Deposits of the new tissue were identified with certainty in six of the cases. In two of the positive cases it was restricted to one and two samples respectively. Where deposits were found, they were always focal, consisting essentially of nodules of the new tissue.

I.B.1. The Liver.

The disease was manifest in this organ in six cases, in one case, P.M. 'D', a large spheroidal tumour in the liver appeared the chief expression of the disease, and in another, P.M. 'I', direct extension into the viscus from neighbouring tumour was the mode of involvement.
In the remaining cases scattered nodules generally small and relatively scanty constituted the form of the disease. The majority of small deposits were in the portal tracts but in one case, P.M. 'O', minute nodules were seen in the lobules.

I.B.2. The adrenal glands.

In only one case, P.M. 'I', was deposit of the growth encountered in the gland. Here there was copious surrounding tumour tissue and the lesion was too advanced to disclose whether the intra-adrenal deposits were primarily intrinsic in the gland or the result of extraneous infiltration. In two cases, P.Ms. 'B' and 'K', glands were invested by Hodgkin tissue but were not the seat of the disease.

I.B.3. The Hypophysis Cerebri.

The gland was not examined in three cases, P.Ms. 'A', 'C', and 'G'. In the remaining ten it was invariably free from deposits.

I.C.1-4.

The lesion was not encountered in the skin nor fascial planes in any case. Further deposits were never found in the omenta other than in organised lymphoid tissue (nodes) in this site. In eight cases a nucleus pulposus was examined since these structures are reputed to represent persistent primitive mesenchyme. The lesion was not present in any.

II. Respiratory System.

In nine cases deposits were present in the lungs. In most it was impossible to surmise where the initium was located. Centrally in the lesions the normal lung tissue was destroyed but peripherally the new tissue appeared to grow out into the alveolar spaces using the alveolar walls.
walls as a scaffolding. The appearances often recalled a chronic interstitial pneumonia. Necrosis was common. In four of these cases nodular deposits were present in the submucosa of the trachea or bronchi. In one case nodules were found in the air passages but not in the lungs.

III. Alimentary System.

In three instances deposits were present in the stomach, in one case a deposit was partly oesophageal as well as gastric. In each case the disease foci presented as small malignant ulcers usually with a few satellite nodules. Persistent but decayed lymphoid tissue was common in this system (para-enteronic lymphoid tissue).

IV. Serous Membranes.

The meninges were not involved in any of the cases where the head was examined. In six cases the pericardium contained deposits and in six the pleura was similarly affected. Peritoneal deposits were observed in four cases.

V. Genito-urinary System.

Renal deposits were encountered in two cases. In another a ureter contained the growth in the submucosa. The bladder was involved in two instances and the prostate gland in one. (All these lesions were mutually unrelated). The ovary contained a deposit in one case.

VI. Cardiovascular System.

Extension from the epicardium into the myocardium was observed in one case.

VII. Skeletal Muscle System.

In two cases infiltration of the diaphragm was present; in one the right psoas muscle was also affected.
VIII. Endocrine Glands. (less hypophysis cerebri and adrenal glands).

In one case the thyroid gland was apparently completely replaced by tumour.

Stability of the lesion.

In the section devoted to the description of typical Hodgkin's disease it was accepted that the microscopical complex generally followed a sequence of maturation in which it retained its identity. While the appearances observed in this process shew variation they have limits. In some of the biopsy series alteration beyond this reasonable latitude was encountered and here further observations on this type of metamorphosis are made. In eleven of the cases appropriate comparison with antecedent biopsy findings can be made and in all the cases the necropsy material itself can be similarly studied.

The decision whether alteration is evidenced or not is dependent on interpretation and judgment. Accordingly there is inevitably an element of arbitrary distinction in these results. The criteria of definition have been applied rigidly; this is reflected in the fact that in the whole series the biopsy of only one case conforms to the ideal typical Hodgkin's disease. The remainder are atypical. In most the atypism is slight and insufficient to assail the diagnoses as practical and truthful reports. In two cases the deviation is more pronounced and in one, the lesion, while simulating cellular tumour-like Hodgkin's disease, must be regarded as belonging to the category of reticulum cell sarcoma.

An analysis of the findings in the series is given in table
<table>
<thead>
<tr>
<th>Case</th>
<th>Biopsy Findings</th>
<th>Necropsy Findings</th>
</tr>
</thead>
<tbody>
<tr>
<td>A.</td>
<td>(None)</td>
<td>(a) Typical mature Hodgkin's disease</td>
</tr>
<tr>
<td></td>
<td>Slightly atypical Hodgkin's disease, eosinophiles reduced, cellularity and pleiomorphism enhanced.</td>
<td>(b) Hodgkin's sarcoma.</td>
</tr>
<tr>
<td>C.</td>
<td>Slightly atypical Hodgkin's disease, eosinophiles low, many foci of epithelioid cells.</td>
<td>Similar to biopsy but eosinophiles further reduced, near to Hodgkin's sarcoma.</td>
</tr>
<tr>
<td>D.</td>
<td>Slightly atypical Hodgkin's disease, mostly small cells, the lesion is early.</td>
<td>(a) Typical Hodgkin's disease.</td>
</tr>
<tr>
<td></td>
<td></td>
<td>(b) Hodgkin's sarcoma.</td>
</tr>
<tr>
<td>E.</td>
<td>Slightly atypical Hodgkin's disease. Eosinophiles few. Focal reticulum cell proliferation.</td>
<td>(a) Atypical Hodgkin's disease, as in biopsy.</td>
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<tr>
<td></td>
<td></td>
<td>(b) Cellular tumour-like Hodgkin's disease, eosinophiles retained.</td>
</tr>
<tr>
<td>F.</td>
<td>(None)</td>
<td>(a) Typical Hodgkin's disease.</td>
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<td></td>
<td></td>
<td>(b) Hodgkin's sarcoma.</td>
</tr>
<tr>
<td>G.</td>
<td>Atypical Hodgkin's disease, no eosinophiles, tumour-like.</td>
<td>Very similar to biopsy.</td>
</tr>
<tr>
<td>I.</td>
<td>(a) Atypical Hodgkin's disease, soft, cellular, eosinophiles low.</td>
<td>(a) Very small component of mature Hodgkin's disease.</td>
</tr>
<tr>
<td></td>
<td>(b) Reticulum cell sarcoma.</td>
<td>(b) Reticulum cell sarcoma.</td>
</tr>
<tr>
<td>M.</td>
<td>Reticulum cell sarcoma, close affinity to cellular Hodgkin's disease.</td>
<td>Very similar to biopsy.</td>
</tr>
</tbody>
</table>
It is seen that in seven of the thirteen cases there was significant alteration in the microscopical appearances of the lesion. In two of these, Cases A. and F. previous biopsy had not been performed but in both the necropsy material displayed at least two reasonably distinct pictures. In these cases the inference is that the Hodgkin's sarcoma form arose on a basis of ordinary Hodgkin's disease since transitions between the two were observed. Of the other five cases shewing alteration of form, three, Cases D., E. and I., shewed similar mixed pictures. In the remaining two cases, C. and H. the alteration was less in degree and incomplete.

In the six cases where metamorphosis was not a feature it is notable that in four, P.Ms. G.J.K. and L., biopsy already revealed tumour-like appearances of the lesion. In one case, B., the atypism was slight. The final case was already frank tumour.

The nature of the metamorphosis.

In general terms this is chiefly expressed by the disproportionate augmentation of morbid reticulum cells in the lesion. This increase affects the morbid mononuclear types, and the multinucleate giant cells. In most cases fibrosis persists but is less in amount than in the classical mature Hodgkin's disease lesion. The eosinophiles diminish to the point of disappearance and frequently the lymphocytes are markedly decreased.

In the earlier part of this work (p. 178) the lesion, which is probably correctly regarded as a morphological variant of the typical form, received the name Hodgkin's sarcoma. These present studies support this view. There is not convincing evidence of any enhancement
of malignancy where this results. On the other hand when the metamorphosis yields a reticulum cell sarcoma in which pleiomorphism is less evident, e.g. case I., truly malignant features are displayed.

The time over which these metamorphoses develop is difficult to assess. In the five cases where the dates of biopsy are known an average period of three-and-a-half years elapsed, but the minimum time was three months and the maximum nine years. In one of the cases, A., specific treatment was not given; all the remainder received X-ray therapy.

The chief effect exerted by X-ray therapy on the lesion is the destruction of lymphocytes and their immediate precursors.

Note on the incidence of tuberculosis.

Histological evidence of tuberculosis was found in four cases. In each, the lungs always shewed foci and in two spread to other organs had resulted. In none was tuberculous disease seriously extensive. Where spread beyond the lungs was observed the lesions were mostly miliary and probably represented terminal dissemination.

Note on Oedema.

Dropsy was very common. In only two cases was it absent. Effusion into the serous sacs affected principally the pleurae, then the peritoneum and least the pericardium. Anasarca was found in nine cases.
PART IV

FIBROSIS AND EOSINOPHILIA

SECTION A.

The phenomenon of fibrosis with observations on its tumour-like character in certain diseases. p.240-262

Illustrations.

Volume III. Figs. 206-212.
Foreword

In the previous part fibrosis and topical eosinophilia in Hodgkin's disease have received attention. It appeared relevant to enquire into these phenomena in the field of general pathology to try and derive information of any comparable findings in other neoplastic or tumour-like diseases.

This has entailed the study of several diseases in which fibrosis appears integral, and an analysis of the incidence of eosinophilia in a series of carcinomata.

Introductory remarks on Fibrosis

Despite wide and common application in Pathology, fibrosis is a process which it is difficult to define comprehensively and accurately. Its etymology does not yield particularly satisfactory understanding. The suffix 'osis' implies an increase of cells, tissues, or stored substances; Powell-White (1927); and, compounded with the root word, fibre, indicates simply an increase of fibres. The exact nature of these fibres is rarely stated fully, but it is generally accepted that they are collagenous, and possibly also fibroglial; Mallory (1913). Mallory also considered that elastic fibres were elaborated by the same cells as the foregoing, but there is reluctance among more recent writers to acknowledge this c.f. Cowdry, (1932), Le Gros Clark (1947). Furthermore, the term elastosis has come into wide use to connote augmentation or undue prominence of these fibres and it is customary to regard them separately. In ordinary reparative fibroblastic activity these fibres
are generally conspicuously absent.

Reflection on the manifestations of the phenomenon of fibrosis regarded as a process immediately disturbs the aptness of the simple definition indicated above. While it seems true in the main to postulate that collagen fibres are a maturation product of proliferating fibrocyte precursors this is not always obvious. In the healing wound the process is patently related to a preliminary proliferation of fibroblasts. In most orthodox teachings it is stated that these cells arise from pre-existing fibrocytes at the locus by an initial de-differentiation followed by homoplastic multiplication of these more primitive cells. These mature as healing proceeds and pari passu fibrosis occurs.

On the other hand the exact mechanism of fibrosis in many of the slowly evolving varieties of replacement fibrosis is less easy to follow. In many examples of ischaemic fibrosis or in arteriosclerosis without hypertonus the sequences in the process are postulated rather than demonstrated. The fibrosis of atrophy, especially in the process of ageing, suggests a retrogressive change by metamorphosis of existing elements rather than by new formation of connective tissue.

Fibrosis is certainly not the exclusive property of the orthodox fibroblasts. Under certain conditions other mesenchymal cells can produce collagen. The lymph node reticulum and homologous tissue elsewhere is quite reasonably and properly regarded as peculiar connective tissue, and as a corollary its fibre product, reticulin, is simply collagen in a special form, or pre-collagen. The difference between collagen fibres and reticulin fibres, frequently made an issue of much importance.
importance is probably over-emphasized. Reticulin increase is demonstrable in certain incipient fibroses but it is not apparently an invariable stage. The fact that reticulum cells can produce collagen is well established and moreover it appears that in some morbid conditions they do so characteristically.

Collagen may arise from non-mesoblastic sources. This phenomenon does not enjoy universal recognition yet, but authoritative opinion in some quarters favours it. Masson considers that the collagen produced in certain tumours or experimentally induced tumour-like growths, of the peripheral nerves is properly attributed to the neurilemmal cells. These cells arise by differentiation from the tissue of the early neural crest and also are reported to emerge from the neural tube by way of the ventral roots, Arey (1941). The exact origin of the neural crest is not agreed (i.e. whether it is simple ectodermal or neurectodermal) but whichever view is more correct it is still fundamentally ectodermal. Masson insists that the only mesoblastic collagen in these growths is that in close relation to the blood vessels.

The application of tissue culture to tumours arising in relation to nerves supports the view that neurilemmal tissue has specific characters and can form collagen, Murray and Stout, (1942). If these and Masson's interpretation are accepted the principle is important, since it enhances the evidence against the older postulates of germinal layer specificity.

There is some evidence that under exceptional circumstances collagen may be formed by endothelial cells e.g. the serosal cells, Cappell, (1952).
The exact relation of that vague but unquestionably valuable entity termed 'hyaline', with collagen, is equivocal. Collagen certainly becomes hyalinised, from a microscopical point of view, and is usually considered a degenerative change; at the same time many other sources of apparently identical hyaline exist. It is very common for example in smooth muscle cells in blood vessels, in the ageing uterus and in the prostate gland. In this particular cell it usually appears as a central thread extending along the long axis of the fibre. In skeletal muscle hyalinisation can produce an appearance identical with collagen.

The current impression of the significance of the phenomenon is that it is compensatory, this term being used in its widest sense. Whilst the untoward effects of its results are widely recognised, the process of fibrosis itself is pretty generally hailed with enthusiasm, much as laudable pus was formerly greeted. It is possible however that too much emphasis has been laid on this rôle. It is continually stressed how important fibrosis is in relation to repair, to inflammation, and how it forms a readily available and cheaply maintained substitute for more specialised tissues. Inevitably one comes to think of it as a simple altruistic process of a benign, plain, work-a-day tissue with no character of its own. The comparative rarity of fibro-sarcoma certainly reflects credit on its general good behaviour.

Nevertheless, the process of fibrosis sometimes deviates from the altruistic path without getting into serious trouble. Little systematic study has been accorded to these biological flutters as a group, although Adami (1911), and Powell-White (1927), offered some tentative outlines.
outlines. An innate perversity to do the wrong thing first probably exists in cells as well as huge collections of cells. Illustrative of this wayward propensity are a few peculiar fibroses which reflect an unwonted degree of enterprise on the part of this rather stodgy tissue. At the same time these growths of fibrous tissue do not receive unanimous approval as examples of neoplasm.

The particular observations offered are admittedly a somewhat quaint collection. They defy any rigid classification, and some indeed are almost anecdotal. However there are reasonable grounds to associate them because in some degree they all shew a process of fibrosis in which some autonymy obtains. This type of fibrosis belongs to that mysterious no-man's-land or borderland dim (Boyd) where the answers are 'well, yes and no'.

**THE PECULIAR FIBROSES**

It is considered that in the following conditions this quasi-neoplastic propensity is evident.

1. Cheloid.
2. Desmoid tumour.
3. Dermatofibroma.
4. Riedel's struma.
5. Polyserositis.
7. Neurofibromatosis.

It is appreciated that these are diverse instances of disease and that many authorities would demur at their consideration as a group. It is not advocated here that they are related beyond evidencing this intrinsic peculiarity in varying degree. In the first four examples the lesion is of local character and in the remainder it is of more general distribution.

/Cheloid
CHELOID

In accordance with Jeigert's ingenious Law - that the degree of reparation of injured tissue always exceeds the demand - fibrosis in the healing process is always more abundant than is really necessary. In the normal case this excess is insignificant, but in some individuals there exists a congenital and sometimes familial tendency for this increment to become embarrassing. This results in abnormal but self-limiting overgrowth of fibrous tissue and constitutes the well-known cheloid or keloid. (The word is variously derived - the former on better etymological grounds). The new fibrous tissue matures rapidly to form large cylindrical deposits of collagen interspersed with rows of nuclei and merges imperceptibly with the adjacent normal connective tissue.

The intrinsic cause of this remarkable predisposition in the subjects of cheloid is obscure, many writers consider that is particularly common in negroes. The tendency becomes apparent after the operation of some form of trauma. This may take many forms, such as burns, surgical wounds, scratches, and etc. and it has been observed after very trivial injuries e.g. vaccination. It is probable that spontaneous development of cheloid is to be related to insignificant traumata operating where the threshold is very low.

In most ordinary texts pronouncements on the interpretation of cheloid are evasive. Authoritative statements from the sources indicated below reflect the diversity of opinion. Adami (1909), included it under the head of proliferative fibrosis, a category in which he placed encapsulating fibrosis, the fibrosis of infective granulomata,
and post-inflammatory fibrosis. Mallory (1913), considered it to be hypertrophied scar, in a category by itself, but he described it in association with fibrous tissue tumour. Ewing (1929) allowed that, 'The neoplastic properties in cheloid are not pronounced'. It seems inferred that it is a neoplasm yet he qualified this implication adroitly by attributing it to chronic disturbance of nutrition. Donaldson (1931) described cheloid as a form of fibrous tumour. Willis (1949), decided in favour of some non-neoplastic disturbance of reparative growth.

Morbid Anatomy of Cheloid

By general definition cheloid is limited to the integument, but homologous formations are observed in bowel, vagina, and possibly other organs.

The naked eye appearance shew considerable variation. Minimal manifestations of the phenomenon are not very rare in striae gravidarum. In the common form encountered in linear wounds, they form elevated flat broad thickenings of the scar which may spread slowly locally, or generally. The overlying skin is often smooth shiny and thin, with loss of elasticity. The colour tends to remain pink longer than normal scar because devascularisation is slower. In some cases, especially after burns, the growths are more exuberant and form large polypoid masses. Several examples of this gross manifestation are impressively illustrated by Mallory (1913) and Ewing (1929).

Microscopically the established cheloid consists of large interwoven bundles of fibres composed of collagen; these are often hyalinised and compacted into a cylindrical form reminiscent of skeletal muscle.

/Elastic
Elastic fibres are not represented. The nuclei are disposed chiefly in the interstices in linear fashion. In form they approximate closely to those of fibroblasts but are larger and often slightly hyperchromic. The periphery of the deposit is characteristically poorly demarcated; it is usually more cellular and merges imperceptibly with the adjacent normal tissue. The blood vessels are well formed and small, but generally numerous. The glands of the dermis and the hair follicles tend to be displaced rather than overrun; the overlying skin is likewise devoid of these structures.

It is submitted that the mild atypism of the components, the purposeless nature of the formation, the tendency to recur and its very limited capacity to infiltrate, warrant its inclusion in the series listed above.

DESMOID TUMOUR

Desmoid tumour is probably the most widely used term to connote the growth which was formerly described as recurrent fibrous tissue tumour of the abdominal wall. The first recorded cases are attributed to Macfarlane (1832) a former surgeon in this Institution, Stewart and Mowat (1924). The name desmoid tumour was introduced by Sänger in 1884, and derives from the supposed similarity of the cut surface to tendon. The main facts regarding these formations are well known and classical treatises have been offered by Pfeiffer in 1904 and Stewart and Mowat twenty years later. Briefly these distinctive and rare growths are recorded chiefly in parous women in the third and fourth decades.

Morbid anatomy of desmoid tumour

The growths are believed to arise from the muscular aponeuroses of
the abdomen, generally that of the m. rectus abdominis. The majority are situated below the umbilicus and almost invariably away from the mid-line. Their naked-eye appearance was well described by Macfarlane (1832). In his first patient the tumour was 'the size of a lemon, greyish, not unlike half-bleached wax, in some parts half transparent, and exhibited a smooth compact texture its centre being fibrous'. Microscopically they resemble fibroma but in some cases their structure gives rise to difficulty in deciding their true character. They may simulate leiomyoma or neurofibroma closely. The growing edge is devoid of a capsule and the overrun skeletal muscle undergoes atrophy. Commonly syncytium-like masses with numerous nuclei are to be seen. This is a well-known phenomenon in damaged muscle and was first fully described by Durante (1902). Adhesions to local intra or extra peritoneal structures are common and probably initiated the belief of Huginer (1860) that they arose from periosteum. This view was long supported by the French. The tendency to recur is possibly over-emphasized and is a relic of the results of less satisfactory surgery. In Stewart and Mowat's series of seven cases recurrence was not recorded but the observation periods were mostly short. In three examples of the lesion studied personally recurrence was not observed. (One of these cases had been treated by irradiation after surgical resection of the main tissue mass).

There is general statement that these growths are tumours; Stewart, Ewing, Willis, Donaldson, Boyd, Muir and many other subscribe to this view. It would be impertinent to question this opinion but the entity deserves some special qualifications. The relation to trauma is remarkably well attested, possibly better than in cheloid. The general
relation of most tumours to trauma is less clear cut. The absence of a capsule is unusual in benign tumours. This of course applies to the established neoplasm, the initium is presumably unencapsulated and the capsule develops in the host tissue in response to the growth of the tumour. At what point this occurs or the factors determining it are matters for speculation. Ewing (1929) obviously favoured the idea that some degree of aggressiveness was essential to wall-building resentment of the host tissue. I have encountered several 'fibromata' of ovary in which the appearances were typical apart from the lack of a capsule, indeed the difficulty of recovering a fibroma suitable for class purposes is notorious.

It is the usual experience that this vigorous fibrous tissue in desmoid does not assume true malignant status. Willis (1949) reported an instance in which one of those growths co-existed with a fibrosarcoma elsewhere and submitted that this might be argued as evidence of a tendency to fibroblastic neoplasia. It is submitted that this growth has some affinities to cheloid but is generally a more aggressive expression of the process.

DERMATOFIBROMA

This peculiar growth has recently enjoyed renewed interest, in fact it is almost true to say that it has been rediscovered.

It is an unquestionably rare lesion and has been recognised under different names over a considerable time.

Among the titles which have been used to designate it are:- sclerosing angioma, fibroxanthosarcoma, benign dermatofibrosarcoma, fibrome-en-pastille, dermatofibroma, histiocytoma and fibromatoid /granuloma.
granuloma. With the exception of the last synonym, which incidentally is said to have been the first name, all the remainder infer etymologically that the growth is a tumour. It is felt that there are grounds for disputing whether this lesion is a true neoplasm and some of these originate from the derivations of the alternative labels which have been suggested to connote it.

The names which acknowledge relationship to angioma cannot imply tumour status on that account. Most present day opinion is hostile to the belief that angioma is more than a malformation, (the very rare angiosarcoma is excepted), accordingly the category hamartoma would be more appropriate. In several further respects this idea has claims to attention. The tumour endothelioma has already received attention (Part I), it will be remembered that while the lucid criticisms of Willis are generally sustained, reasons for sympathy with it were offered. Amongst these was the submission that the tendency to form whorls could be interpreted as a property beyond simple functional modification. It could be regarded as an atavistic trait of endothelial cells. In the few examples of dermatofibroma which I have encountered at first-hand this tendency has been discernible in some degree. The production of collagen by endothelial cells of capillaries has been demonstrated by Corner (1920). In most dermatofibromata sclerosis is a prominent feature but vaso-formative habit is far less in evidence so that sclerosing angioma is a far-fetched nomenclature for the growth.

Xanthoma is a neoplasm which has come down in the world. The name frequently appears in inverted commas, a slight on its individuality, or it is joined with another or several other syllables giving high-sounding
compound titles which are no longer popular. It has an extensive and confusing literature in the older dermatological annals and in the modern oncological works it receives scant attention. The inference gained from the latter is that it is a very dubious neoplasm. That aggregates of xanthoma cells appear in certain tumours is widely known but there is always hesitation to ascribe neoplastic properties to them.

The view that the growth is a variety of fibroma or benign sarcoma of fibrous tissue has certain claims. An unencapsulated fibroma of ovary is widely recognised and has been alluded to earlier in this section; the other alternative, benign sarcoma while it is paradoxical doubtless springs from recognition of this anomaly.

Woringer's term, histiocytoma, (1931) is probably the most popular name at present. This title was prompted by the observation that cells of the growth store iron and lipoid. It has been supported by the experimental finding that cells of the growth could ingest and store colloidal iron injected into the tissues in their vicinity. Since the growth scleroses and fibrifies as it matures it follows that histiocytes (macrophages) form collagen. This is an undoubted fact in experimental work, Curran, (1951), Nicol and Abou-Zirky, (1953), but most human morbid anatomy texts are hesitant to postulate the process with assurance in natural disease. The principle is very reasonable, in a healing wound the distinction between fibroblasts and macrophages can be so difficult that their identity is the only logical interpretation.

An interesting corollary of accepting Woringer's hypothesis is that we are furnished with an example of benign tumour of the reticuloendothelial system. The rarity of these neoplasms has already been
alluded to in Part I. where the claims of myxoma were upheld. Further to this exceptional status is added the fact that the deposit is almost always solitary. In the rare histiocytic reticulosis (of both sinus and medullary type, Robb-Smith, 1938) systematization or at least very wide diffusion was the key-note.

The name fibromatoid granuloma is credited with being the oldest. Except in the sense in which Ewing employs the word, granuloma is an unsatisfactory term, like tuberculoma or syphiloma its employment is chiefly sanctioned by usage. As ordinarily understood it assumes an inflammatory process. The lesion in question here does not suggest inflammation if only on account of the total absence of chronic vascular dilatation.

Morbid anatomy of dermatofibroma

The lesion is invariably benign, it is nearly always solitary, of very indolent growth and although often circumscribed, it may be diffuse. In the fresh specimen the intracutaneous site, the remarkable induration, the absence of capsule and the faint flat yellowish-grey colour of the cut surface are characteristic. The exact nature of the colouration is obscure, it is unlikely that haemosiderin accounts for it. Microscopically it generally presents as an imperfectly circumscribed proliferation of cells. The bulk of these has that superficial appearance of uniformity which is common in tumour. This moiety is generally composed of stout spindle-shaped cells closely resembling hyperchromic fibroblasts, frequently the nuclei shew distortion and the cells then appear stellate. They are applied to or set in a mesh of collagen fibres. In this groundwork tissue and at the periphery xanthoma cells
may be identified either singly or in groups. A variety of giant cell
the Touton cell (1885) is commonly present, this is believed to be a
fusion product of xanthoma cells. Haemosiderin is nearly always demonstrable in some areas of the growth.

It is important to appreciate that much of the theory regarding this
tumour-like formation is speculative; the precise histogenesis has not
yet been satisfactorily proved. The synovioma or benign giant-cell
tumour of tendon sheath presents a fairly close similarity in structure
to dermatofibroma but its claims to tumour status are much more justified.

In summary while the growth does present some tumour-like propensities its pretensions in this respect are low, and it may be regarded as a link between the peculiar fibrosis under discussion and neoplasia.

**RIEDEL'S STRUMA**

This uncommon malady furnishes another example of localised peculiar fibrosis. Unfortunately there does not appear to be any concise information on exactly what constitutes the disease nor is there agreement on its nature, aetiology, or natural history. There are, nevertheless, sufficient premises to consider it a separate disease entity but with the qualification that it is still imperfectly defined.

In 1896 Riedel described a form of chronic inflammation of the thyroid gland which transformed it into a bulky tumour composed of dense fibrous and sclerotic tissue. The entire gland was involved and converted into a mass of almost iron-hard consistency, and so generally adherent that extirpation was difficult or impossible. He made no mention of extensive lymphocytic infiltration nor the formation of germinal
germinal follicles in the gland. He further observed that the disease did not behave like carcinoma, for which it was commonly mistaken, and that it did not respond to appropriate therapy for the common specific chronic inflammations. Since this initial communication the definition has been widened by some observers, and narrowed by others with the result that misunderstandings have been caused. The characterisation of rare diseases is always a matter of difficulty for the same reason that ex pede Herculanem is an unsound principle for a physician. Not a few rare conditions have been described on very small series, e.g. the uncommon nervous disorders, endocrine gland dysfunction syndromes, and etc., and it is inevitable that errors are made.

Mindful of the succinct injunction of Dr. Ferguson-Smith, - which same he was wont to repeat with enviable precision and rapidity 'The common forms of rare diseases are rarer than the rare forms of common diseases!' a certain latitude in definition is surely justified and this does not reflect criticism on the original observations.

The nature of the disease is unknown but this of course does not preclude copious speculation which is well seen in the numerous titles suggested; sic: Ligneous thyroiditis, woody thyroid, primary chronic inflammation of the thyroid gland, Riedel's thyroiditis, iron-hard tumour of the thyroid gland, and benign granuloma of the thyroid gland, all reflect the varying ideas. Riedel's struma is tactfully non-committal. Ewing, who suggested the penultimate name in the list, clearly conceived of 'a granuloma of neoplastic type', the idea pervades his unquestionably solid contributions to oncology, this name possibly expressed a belief that the disease has tumour-like qualities.

The aetiology is also unknown. The relation of this disease to
/Hashimoto's
Hashimoto's disease is still a vexed question some observers maintain they are distinct e.g. Hellwig, (1938) Boyd, (1945); whilst others affirm that they are simply stages in a single process Shaw & Smith, (1925). The former evidence is based on the retention of the same microscopical picture in serial biopsies made over protracted periods and the latter on studies of quantities of material in which gradations from one to the other can be traced. Most orthodox opinion at present favours the former hypothesis.

Boyd (1945) confesses his own bewilderment at the 'mixed' types one meets in actual practice. In this connection it is quite probable that 'burned-out' Graves' disease and myxoedema supply a number of these equivocal cases.

The natural history of the lesion is elusive, because so frequently it is fully established when the first examination is made. It is debatable whether studies of large series shewing progressive degrees of fibrosis are admissible as evidence of the process particularly in a structure like the thyroid gland. It has been suggested, De Courcy (1943), that the lesion is primarily a peri-adenitis but the morbid anatomy points rather to an augmentation of existing fibrous tissue in the stroma of the gland as well. This proceeds centripitally bearing down on the acinar tissue and at the same time reducing its blood supply. This mechanical crushing and starvation result in atrophy of the epithelial moiety. At the same time fibrous tissue expands outwards driving contiguous lobules of the gland further apart. In this way absolute increase in size of the gland is attained.

The main interest here is the fibrosis which fixes the gland so firmly.
firmly to adjacent structures. This has more than a passing resemblance to desmoid tumour and is in fact closely parallel. Fixation is probably an essential feature in the disease. In a series of 23 cases collected by Shaw & Smith (1925) 18 shewed this phenomenon, in the remainder the records were incomplete. In a series observed personally the phenomenon was present in 3 cases out of 4. The infiltrative tendency of the fibrosis is much the same as in desmoid.

POLYSEROSITIS

This term is used to include a group of rare conditions in which there is remarkable fibrous thickening and sclerosis of the serous membranes, with a tendency to chronic dropsical effusions into serous sacs. Many other names have been attached to the disease and several to conditions which are regarded as incomplete manifestations of it. The term pervisceritis connotes the same disease in France; in Italy, Concato's disease is identified with the condition and this eponymic title is frequently used in the United Kingdom. Formerly polyorrmenitis chronica was employed synonymously in parts of Germany. A frequently quoted authority, Kelly (1903), used the name 'multiple serositis'. It is probable, Cappell, (1951) that Picks disease (pericarditische pseudolebercirrhose, 1896) is of the same essential nature.

This multiple nomenclature and the general vagueness of the subject have given rise to extensive searches of the literature to ascertain who was the original sponsor. In the publications available to me many of the early references are to inaccessible sources, chiefly continental in origin. Kelly considered that the first case was reported by van Deen in 1846 but other authorities attribute priority to Lancisi (1728),
Corvisart (1812), or Chevers (1842).

In the third edition of Corvisart's treatise (1818) there is an account of a case of chronic constrictive pericarditis in a man aged 40 years. The case notes and post mortem findings indicate that this was probably an example of polyserositis. Two other cases of adherent pericardium are described in this section but their identity is dubious.

Chevers described two cases of chronic constrictive pericarditis with ascites, in the first of these (Case Jane.P.) a pleurisy was also present and was doubtfully ascribed to a chronic empyema.

Kelly (1903) published an instructive review of the condition including a full account of a fatal case which he examined himself and 39 cases extracted from the literature. He was the first author to regard Pick's disease, chronic constrictive pericarditis, chronic hyperplastic peritonitis & etc., as being merely different expressions of the same morbid process, and his paper has formed the basis for subsequent work.

Interest in the condition was much stimulated when surgical treatment was found to give considerable relief or even cure in a high proportion of cases. White (1935) related a series of 15 personally observed cases, some of whom were successfully treated and Burwell & Blalock (1938) a further group of 19 cases. The aetiology of polyserositis is not agreed. In Kelly's case careful microscopy failed to reveal any evidence of tuberculosis. Four out of the 39 abstracted cases were considered tuberculous in origin. In White's series the causation was ascribed to tuberculosis in only 2 cases with certainty. White insisted that Pick's disease was not related to polyserositis. In Burwell & Blalock's patients 16 out of the 19 were considered of tuberculous origin,
origin, 3 due to pyogenic microbes.

The morbid anatomy is variable depending on the extent of the disease. The process is essentially a remarkable hyaline thickening over the affected sites. This is chiefly collagenous white fibrous tissue with sparse cellular content and of very low vascularity. In some cases organs invested by these laminated deposits show extension of fibrosis through their capsules suggesting a feeble infiltrative tendency at work. So far as can be understood the process appears to be naturally slowly progressive. While it has been thought that the lesion might owe its character to some chronic mild irritant this has not been well proved. It appears reasonable to regard it as an example of peculiar fibrosis since it persists in the absence of any identified cause in most cases, and its wayward if slow progress reflects a degree of autonymity.

**MYELOFIBROSIS**

This is a rare condition in which the haemopoietic elements of the bone marrow are replaced by fibrous tissue. The replacement appears to be a slowly progressive process which ultimately involves all the marrow previously occupied by active blood-forming tissue. Pari passu extra-medullary foci of haematopoietic tissue develop especially in the liver, spleen, and the lymph nodes. Small foci have also been observed in the kidney, pancreas, abdominal fat & etc.

In common with all rare disease (under 200 cases had been reported in 1946) the characterisation is imperfect and there is some doubt as to whether it really is a morbid entity.

On the other hand it is reasonable to suggest that myelofibrosis is a process
a process which is possibly fundamental in character and recurs as a common factor in certain specific diseases and syndromes.

The phenomenon has been recognised for a long time. Most authorities attribute the initial observations to Henck in 1879. This worker described two cases. One of them was an instance of lymphatic leukaemia in which fibrosis of the bone marrow was found at necropsy.

In many respects the feature of myelofibrosis has been overshadowed by the results related to it. Thus after the description of leucoerythroblastic anaemia by Vaughan, (1936), and the conception of 'myelophthisic' anaemia, this blood picture has dominated the subject. While a pure primary myelofibrosis certainly exists and does characteristically produce this anaemia many other lesions can bring about the same blood change and numerically they are overwhelmingly more frequent. Metastatic carcinoma in bone marrow is probably the commonest cause of this; most instances are recorded in cases of mammary, prostatic, and pulmonary carcinoma.

Confusing the issue further are a series of very rare diseases and syndromes. The marble-bone disease of Albers-Schönberg and Albright's syndrome are the best known examples, and the storage diseases of the reticulo-endothelial system including Gaucher's disease, Niemann-Pick's disease, and Schuller-Christian's disease may also give rise to this anaemia.

Unfortunately there is not any authoritative identification of a large number of doubtfully characterised conditions related to these. Albright's original name 'osteitis fibrosa disseminata' - which lesion is mainly unilateral - would seem the same as Lichtenstein and Jaffe's /polyostotic
'polyostotic fibrous dysplasia' in so far as the bones are concerned. The term osteosclerosis is sometimes used instead of myelofibrosis. Mettler and Rusk (1937) use it to include osteopetrosis (Albers-Schönberg's disease) and myelofibrosis. Myelosclerosis is apparently employed synonymously with myelofibrosis.

Finally there is a chaotic group of cases under the headings of 'leukaemoid blood picture', aleukaemic myelosis with osteosclerosis (Stevens and Bredeck, 1933), and unusual cases of polycythaemia vera.

This subject leads into that darkness which stultifies the most diligent efforts to understand it properly. There is however one paper which is most helpful. This is the communication of Vaughan and Harrison (1939). These authors suggest that the underlying cause is a 'progressive hyperplasia' of multipotential mesenchymal tissues by an unknown stimulus. The leuco-erythroblastic activity they believe may be partly secondary to the myelosclerosis and the fibrosis itself may in part be the result of the haemic change. This idea is the reticulosis idea, but quasi-neoplastic fibrosis is surely an equally reasonable name for the process.

NEUROFIBROMATOSIS

This subject invites controversy from the start. An uncommon disease, yet widely known about, it has evoked violent argument since its initial description. Von Recklinghausen (1882) first emphasized the relation of the lesions to the peripheral nerves and his name is now commonly attached to the condition, the disease was nevertheless well known at least twenty years prior to this, (Hitchcock 1862).

Opinion has backed and veered irresolutely with regard to its
nature, status, and histogenesis for so long that one wonders how long it is to enjoy its present position.

Some few facts regarding it appear generally accepted. The complex is an entity, this despite the protean morphology it may present. Thus the full picture with multiple molluscum fibrosum lesions, pigmentation, plexiform neuroma, hypertrichosis, bone changes, and elephantiasis arabum, represents the entire realisation of the same process as the 'neurofibroma' of skin. The present view is that the disease represents a maldevelopment of the peripheral nervous system, Stout (1949), Willis (1949). For this reason it is proper to term it hamartoma.

The origin of the growth is less easily settled. It is now considered that it is predominantly neurilemmal with admixture of neurites. These latter are conspicuously absent in most encapsulated neurilemmomata. Stout (1949) laid considerable stress on the specificity of neurilemmal cells being established by tissue culture but curiously enough in this account cultures from neurofibromatosis were not recorded. This author pointed out that malignant schwannoma may develop on a basis of neurofibromatosis and offered tissue culture evidence to support the contention that these malignant growths were composed of neurilemmal cells. This however is only indirect evidence that neurofibromatosis is neurilemmal in origin, and it is pertinent that neurilemmoma is almost invariably encapsulated and very rarely goes malignant.

The older views which were well expressed by Ewing have much to support them. If cell morphology is reliable than fibroblasts and endothelial cells are certainly present in many neurofibromata and a
tendency to whorling - as in meningioma is often distinct.

Whether the growth is entirely neurilemmal or partly fibroblastic there is consent that collagen is formed and this is equivalent to acknowledging that fibrosis is a sequel. This fibrosis and simultaneous overgrowth of tumour-like cells whence it is arising may proceed very rapidly. Instances are recorded of relatively sudden pronounced growth of this kind supervening in plexiform neuromata, the process extending up to and even entering the spinal cord, without true sarcomatous transformation of the tissue involved. In a similar fashion a localised elephantiasis neuromatosa may spread from an initially restricted site to involve large tracts of tissue.

Although neurofibromatosis is a less satisfactory example than others which are cited it is submitted that there is justification for including it since certain of its expressions illustrate tumour-like fibrosis.
SECTION B.

Eosinophilia, with special reference to the phenomenon in tumours. p. 264-270

Illustrations

Volume III. Fig. 213

Volume IV. p. 30-33.
Eosinophile leucocytes are found in the blood and certain fixed tissues. In the circulating blood of normal man these conspicuous cells account for 1-4 per cent of the total leucocytes the absolute figures being 50-400 per cu. mm. In the fixed tissue of the adult its normal habitat is the bone marrow, but it is also usually found in the stomach, duodenum, caecum and vermiform appendix. In the foetus and neonatal infant it resides in the archemyelon, viz. spleen, liver, thymus gland, pancreas, and less constantly in other sites.

The disparity in morphology between the blood and tissue types is chiefly attributable to the results of the different preparation techniques conventionally adopted, but eosinophile cells peculiar to extra-myeloid sites also exist.

The morphology and tinctorial properties of this cell are well-known. The nature of the oxyphilic granules is not fully understood. They are insoluble in fat solvents, weak acids, and alkalis. They are negative to the xanthoproteic reaction with nitric acid, and they contain ionisable iron. This last property was first demonstrated by Barker (1894). Despite the negative reaction with nitric acid they have been considered to be protein products since Ehrlich first suggested that they corresponded to the aleurone granules of ova. (Peyton Rous, 1908, Jolly 1922).

The function of the cell is obscure. As a leucocyte it is unimpressive; it does not respond proportionately to the chemotactic stimuli which excite the neutrophiles, and in fact is characteristically suppressed in the acute inflammatory exudate.

The origin of the eosinophile is not universally agreed. Ehrlich held the view that production was exclusively myeloid and this has been /upheld
upheld by Ringoen (1921) and Biggart (1932). This theory has been widened by the inclusion of an origin from extramedullary myeloid tissue, which locus may be the result of myeloid metaplasia, Lang (1926). The mechanism of origin in this phenomenon has been interpreted differently. Dominici (1909) traced the development of eosinophile leucocytes from lymphocytes and Weidemisch (1910) maintained that this represented a process of phagocytosis, the ingested particles being broken down erythrocytes. C.H. Bunting (unpublished observations) came to similar conclusions but considered that plasma cells were the unfed potential eosinophiles. Fallacies attended the early experimental work of this school, and it is now widely abandoned that the granules are derived from effete red cells. On the other hand the transformation of lymphocytes, plasma cells, stem-cells & etc. to eosinophiles is still supported. Pullinger (1932), Gillman et al (1949). Up till very recently it has been singularly difficult to photograph eosinophiles satisfactorily, and many pathologists are sceptical about the objectivity of drawings. This fact has militated against general acceptance of this theory. All considered it appears probable that myeloid tissue produces most of the eosinophiles but that cytometaplasia also contributes some. It would appear reasonable to anticipate an hyperplasia of the precursors of this cell in the myeloid tissue when eosinophilia is present. This however is difficult to substantiate because marrow is impossible to sample accurately and the augmentation may be very slight, Beattie and Dickson (1943). One corollary of the exclusively myeloid origin theory is that topical eosinophilia must be infiltrative; this postulates the presence of some attracting stimulus resident in the tissue.
One further theory is mentioned. Gutig (1907), Downey (1914) held that the blood and tissue eosinophiles were different in every way, and should be considered separate cells. The identity of this tissue eosinophile is distinctive. It is larger than the blood counterpart, and the granules are less distinct and brownish-pink when stained with eosin. These cells are commonly found in lymph nodes and are almost certainly macrophages.

Eosinophilia in the blood has been extensively studied and the conditions in which it occurs have been well defined, Muir (1941), Whitby and Britton (1942). This has encouraged several generalisations of value, but the phenomenon can be capricious in the face of known causes and idiopathic examples exist. There appears ample evidence that certain exogenous proteins precipitate eosinophilia. Endogenous proteins are possibly also capable of producing the effect. Certain drugs, particularly camphor and nervanol, tend to produce eosinophilia, and less constantly mercury and iodoform are credited with the same property.

Local eosinophilia in the fixed tissues has not received the same amount of study as the more readily examined blood. The phenomenon accompanies many cases of blood eosinophilia but these examples can be set aside. There are conditions in which local eosinophilia may occur as an isolated finding, e.g. eosinophilic granuloma, non-specific peri-appendical granuloma, Musgrove and Docherty (1950); ligneous perityphlitis, Wilson, Dockerty, Waugh and Bargen (1949); and periarteritis nodosa, to mention only a few. It is quite common in gastric ulcers, nasal polyps, and gummata.

In sterile abscesses produced by turpentine eosinophilia may be /intense,
intense, and this has suggested necrosis with endogenous protein release as the exciting cause; a similar idea has been offered to explain the phenomenon in X-ray and radium burns.

Eosinophilia in tumours has not been much studied. Mallory (1913) emphasized that the phenomenon was common in carcinoma of the uterine cervix. Biggart (1932) recorded local eosinophilia in all of a series of 19 tumours - 14 malignant and 5 benign recovered from a variety of sites. Willis (1947) discussing the local eosinophilia of Hodgkin's disease considered that 'the granulocytes and plasma cells in Hodgkin's appear clearly to be only reactionary elements similar to those seen in many other kinds of tumour'.

It was considered that information on this problem might aid the interpretation of the local eosinophilia of Hodgkin's disease.

Material. A series of 500 separate carcinomata encountered consecutively in the Surgical Specimen Routine were examined for local eosinophilia. Since lymphoid tissue is mesodermal comparison with sarcomata would have been more apt but the numbers available were far too small.

The tumours were derived as follows:-

<table>
<thead>
<tr>
<th>Site</th>
<th>Total Number</th>
</tr>
</thead>
<tbody>
<tr>
<td>Integument and its derivatives.</td>
<td>213.</td>
</tr>
<tr>
<td>Gastro-intestinal tract.</td>
<td>139.</td>
</tr>
<tr>
<td>Genito-urinary system.</td>
<td>103.</td>
</tr>
<tr>
<td>Respiratory system.</td>
<td>9.</td>
</tr>
<tr>
<td>Miscellaneous (includes lymph nodes)</td>
<td>36.</td>
</tr>
<tr>
<td></td>
<td><strong>Total 500</strong></td>
</tr>
</tbody>
</table>

The bias in favour of tissues and organs amenable to surgery is /manifest,
manifest, the series is not representative of cancer incidence in general.

Methods.

The routine sections stained with haematoxylin and eosin were examined first. Where eosinophiles were apparent confirmation was usually made by staining with dilute eosin or by the carbol-chromotrope method. The concentrations encountered were graded as sparse, +, ++, and ++++. The index ++, approximated to the degree of eosinophilia encountered in typical Hodgkin's disease, namely obvious without search. For the purposes of this survey examples exhibiting ++ or +++ were considered topical eosinophilia.

Results.

<table>
<thead>
<tr>
<th>Total Tumours</th>
<th>Local Eosinophilia</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>500.</td>
<td>75.</td>
<td>15.</td>
</tr>
</tbody>
</table>

Analysis.

1. Integument and its derivatives.

<table>
<thead>
<tr>
<th>Site</th>
<th>Total cases</th>
<th>Eosinophilia</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Skin</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Squamous cell cancer</td>
<td>38.</td>
<td>6.</td>
<td>15.8</td>
</tr>
<tr>
<td>Basal cell cancer</td>
<td>30.</td>
<td>0.</td>
<td>0.</td>
</tr>
<tr>
<td>Malignant melanoma</td>
<td>12.</td>
<td>0.</td>
<td>0.</td>
</tr>
<tr>
<td>Metastatic</td>
<td>5.</td>
<td>0.</td>
<td>0.</td>
</tr>
<tr>
<td>Breast</td>
<td>128.</td>
<td>3.</td>
<td>2.3</td>
</tr>
</tbody>
</table>

Notes. In 3 cases of squamous cell cancer shewing eosinophilia, genital area skin was affected; the total of epitheliomata in this region was 7. All the positive breast cases were spheroidal cell tumours, 1 shewed /Paget's
Paget's intra-epidermal carcinoma.

2. Gastro-intestinal tract.

<table>
<thead>
<tr>
<th>Site</th>
<th>Total cases</th>
<th>Eosinophilia</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mouth</td>
<td>23</td>
<td>1</td>
<td>4.3</td>
</tr>
<tr>
<td>Oesophagus</td>
<td>7</td>
<td>2</td>
<td>28.3</td>
</tr>
<tr>
<td>Stomach</td>
<td>24</td>
<td>15</td>
<td>62.5</td>
</tr>
<tr>
<td>Caecum</td>
<td>4</td>
<td>4</td>
<td>100.0</td>
</tr>
<tr>
<td>Colon</td>
<td>37</td>
<td>13</td>
<td>35.1</td>
</tr>
<tr>
<td>Rectum</td>
<td>20</td>
<td>11</td>
<td>55.0</td>
</tr>
<tr>
<td>Vermiform appendix</td>
<td>2</td>
<td>1</td>
<td>50.0</td>
</tr>
<tr>
<td>Bile ducts</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Gall bladder</td>
<td>3</td>
<td>1</td>
<td>33.3</td>
</tr>
<tr>
<td>Omentum</td>
<td>12</td>
<td>1</td>
<td>8.3</td>
</tr>
<tr>
<td>Liver</td>
<td>3</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Pancreas</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Parotid gland</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Notes. The high incidence of eosinophilia in gastric, caecal, colonic and rectal tumours is probably related to the normal presence of eosinophiles at these sites. Little significance attaches to the other relatively high figures since the total numbers are so small.

3. Genito-Urinary system.

<table>
<thead>
<tr>
<th>Site</th>
<th>Total cases</th>
<th>Eosinophilia</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Kidney</td>
<td>10</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Bladder</td>
<td>18</td>
<td>7</td>
<td>38.8</td>
</tr>
<tr>
<td>Urethra</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Uterus</td>
<td>11</td>
<td>2</td>
<td>18.2</td>
</tr>
<tr>
<td>Cervix</td>
<td>23</td>
<td>3</td>
<td>13.0</td>
</tr>
<tr>
<td>Ovary</td>
<td>13</td>
<td>1</td>
<td>7.7</td>
</tr>
<tr>
<td>Prostate gland</td>
<td>24</td>
<td>2</td>
<td>8.3</td>
</tr>
<tr>
<td>Testis</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

Notes. The figure for bladder cancers is high, most of these growths were papillary carcinomata. The uterine and cervical incidence of eosinophilia is in keeping with general experience. The single positive ovarian
ovarian eosinophilia was in a malignant teratoma from a child.

4. **Respiratory system.**

<table>
<thead>
<tr>
<th>Site</th>
<th>Total cases</th>
<th>Eosinophilia</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Larynx.</td>
<td>6</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Branchial cleft.</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Bronchus.</td>
<td>2</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

**Note.** The series is too small for reliable results.

5. **Miscellaneous.**

<table>
<thead>
<tr>
<th>Site</th>
<th>Total cases</th>
<th>Eosinophilia</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td>Lymph nodes.</td>
<td>30</td>
<td>2</td>
<td>6.6</td>
</tr>
<tr>
<td>Thyroid gland.</td>
<td>4</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Fascia.</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Adrenal (secondary)</td>
<td>1</td>
<td>0</td>
<td>0</td>
</tr>
</tbody>
</table>

**Note.** One of the cases of lymph node metastatic carcinoma was a gastric node, the other was from the scalp.

The limitations of the series preclude authoritative interpretation of eosinophilia in carcinomata generally, but there is evidence that the phenomenon is related to the site of the growth and that it is not so rare as might be expected. It is not related to necrosis but is often accompanied by plasma cell infiltration.
PART V

EXPERIMENTAL

An attempt to reproduce Hodgkin's disease in rats and mice by trypan blue poisoning. p. 272-287

Illustrations.

Volume III Figs. 214-235.

Volume IV p. 89-99
EXPERIMENTAL WORK ON HODGKIN'S DISEASE

Foreword. The experimental reproduction of a disease in a suitable laboratory animal integrates knowledge in a unique fashion. The advantages include simplification and control of conditions beyond what is feasible in natural disease, and curtailment of the duration of observations.

Few maladies have resisted the experimental method with the success of Hodgkin's disease. It has stultified all attempts. The facts attending this recalcitrance deserve emphasis.

Hodgkin's disease is almost certainly confined to Homo sapiens. Claims of its occurrence in lower animals are rare and its identity in these reports is equivocal. MacMahon (1934), reported a case in a dog. The disease was limited to a single fused cervical node mass on one side; the microscopical picture did not conform to the accepted one closely. Stalker, Schlotthauer, and Feldman (1936), recounted another instance in a dog. The lesions were relatively focal and circumscribed; microscopically they were close to cellular Hodgkin's disease. It is noteworthy that these authors interpreted their findings critically and with caution. Medlar and Sasano, (1937) gave an account of supposed Hodgkin's disease in an elderly female rabbit. The photographs of the lesions in this case were not convincing, the fields shewn were very small. Forbus and Davis (1946), reported a reticulo-endothelial system lesion encountered in seven pigs, which they regarded as close to the human disease. This affinity is not borne out by critical analysis of their findings. These reports comprise most of the better substantiated examples, there has perhaps been too sanguine an interpretation of their significance.
significance by some reviewers. Among the older writings Hodgson (1903), and McFadzean (1903), gave brief references to the disease in horses but the evidence was doubtful.

Transmission of the disease to lower animals has never been successfully accomplished. Attempts have been made since Longcope's pioneer experiments (1903, 1907) and many different animals have been used. The general experience is that the grafts die out; if large they may become encapsulated, and if small they are absorbed. Transient reactive hyperplasia of the host lymph nodes has been described in some of the experiments. The universal failure is partly referable to the expected fate of transplants to heterologous species and indicates that an agent capable of inducing the lesion is probably not present in the new tissue.

Tissue culture studies have yielded information about the reticulum cells of the lesion, Grand (1944, 1949), but have contributed little else.

Empirical research has naturally a claim, however irrational it may at first appear. Before more particular account is given, one observation is offered.

It is insisted that the production of an ephemeral lesion having some or even much microscopical similarity to Hodgkin's disease is a very restricted accomplishment. Failure to appreciate this has been a regrettable shortcoming among certain experimental workers. Any student of the human disease cannot fail to be impressed by the relentless if temporarily hesitant progress which Hodgkin's disease invariably presents. This aspect of its behaviour has never been simulated. It is possible to produce lesions with close morphological

/affinities
affinities to the disease by a considerable variety of methods in the common laboratory rodentia. These however are ephemeral, and fail to evolve like the true disease. Examples of this type of reaction are seen in the work of Miller and Turner (1943), who injected elaborately prepared extracts of urine from patients with leukaemia and Hodgkin's disease into guinea-pigs. In this work the fractions used were one containing hydroxy-acids which appeared to stimulate lymphopoiesis, and another made up chiefly of non-carbinol-acids which had a corresponding action on myeloid tissue. It is pertinent that these extracts were deleterious, the animals seldom survived beyond three to five weeks. Very similar reactions could be induced by extracts of beef. The lesions found were well illustrated, but the fields depicted were very small. Initially promising this line of research appears to have been abandoned.

In 1948 Gillman and Gillman employed parenteral injection of trypan blue in female rats in a study from which they drew analogies between the results of this empirical procedure and those observed in babies whose mothers contracted rubella in early pregnancy. In the course of this work they discovered that tumours of reticulum cell type could be induced by this means in several different organs. In the following year Gillman, Gillman, Gilbert and Spence (1949) produced a comprehensive study of these tumours. In addition they reported changes in lymph nodes of trypan blue treated rats which approximated closely in morphology to Hodgkin's disease and Hodgkin's sarcoma. They interpreted these blastomatoid formations as similar to those in man which Cazal (1942, 1946) regarded as manifestations of 'La reticulose histiomonocytaire'. 
cytaire*. In brief the underlying idea of this was an initial reactive hyperplasia which by accentuation ultimately assumed tumour status. Emanating from South Africa, this work naturally recalls Pliny's observation - 'ex Africa semper aliquid novi', and to the sceptical, the General plan of experimental work undertaken. A repetition of the Gillmans' work with trypan blue was attempted. Several modifications were introduced. These included the use of a different brand of trypan blue, its administration to mice as well as rats and alteration in the frequency of injection. A further series of rats were given another vital stain of comparable chemical structure.

As the result of findings established in the preliminary studies, the development of the lesions was followed by serial biopsies in a further series of identically treated rats. The effects of cessation of the trypan blue injections was observed, and attempts were made to transplant the growths obtained.

Material and methods.

Animals.

Male albino rats of an inbred laboratory colony of wistar strain were used. The mice were females which had matured beyond their usefulness for the Ascheim-Zondek test, (these animals were supplied weekly to the laboratory by Tuck and Sons).

They were distributed as follows:-
Series (A. 10 rats, av. wt. 58 gm.
B. 10 rats, av. wt. 56 gm.
D. 9 rats, av. wt. 100 gm.
E. 10 rats, av. wt. 125 gm. Total 38 rats.
C. 15 rats, av. wt. 135 gm. Total 15 rats.
F. 20 mice, av. wt. 16 gm. Total 20 mice.

(Controls. 10 rats, av. wt. 58 gm.
(Controls. 10 mice, av. wt. 16 gm.

Note. A female rat was inadvertently included in Series B. This was overlooked till a litter of eleven was born nine weeks after the experiment started. Nine of the offspring were males and constitute Series D. (The mother was withdrawn from the experiment).

Diet. Rate cake, Rowat Research Institute Diet No. 86. Supplemented with bread crusts, occasional cooked bones, and cabbage. Water ad lib.

Dyes used.

These were made up freshly in sterile distilled water, 1 gm./100 ml.

Injections.
1. Trypan blue. Series A. and B. 1 ml. (= 10 mgm. dye) by intra-peritoneal injection every fortnight. Series D. and E. the same dosage but given weekly. Series F. 0.1 ml. (= 1 mgm. dye) by intra-peritoneal injection weekly.
In Series A., B., C., D. and F., the endeavour was to maintain the animals alive as long as possible. They were killed only if they became moribund; or if they survived, to the limit of the experiment.

In Series E. animals were killed at 70, 100 and 120 days, after which injections ceased. The remaining seven were subjected to biopsy at 120 days; of these three were killed later, at 140, 160 and 180 days (one was found dead at 167 days). The last three had biopsy repeated at 180 days and were killed at 360 days.

Results.

Trypan blue, Series A., B., D. and E.

In this group of 38 rats, 5 were lost due to escapes, cannibalism and mistaken identity.

During life the following changes were recorded. After injection of the trypan blue the animals tended to remain subdued for several hours. Initially they put on weight more rapidly than the controls but after about two months the position was reversed. Generally by the second injection, whether at 14 days or 7 days interval, general staining of the tissues appeared. This coincided precisely with the full description given by Cappell (1929); it did not augment appreciably thereafter. After 2-3 months most of the beasts became emaciated and weak; they sat hunched up and shivering. In many the abdomen became swollen, and their movements ataxic. Anaemia of the order 3 X 10⁶ erythrocytes per cu.mm. was observed. Terminally they sickened rapidly, diarrhoea was common.

At death the common findings were:

1. **Ascites.** The fluid was faintly dye tinged and had a peculiar foul
metallic odour. It was occasionally turbid and infected.

2. **Liver.** The organ was usually considerably enlarged (average for series A., B. and D. 32.3 gm., average for controls 12.5 gm.) It was dark water-chocolate colour, the surface was sometimes very faintly stippled but hardly granular. Tumour-like nodules 1-10 mm. were found in 18 cases (54 per cent). These varied in colour from pale sky-blue to deep royal blue. They were scattered indiscriminately through the liver substance. Central necrosis was commonly visible in the nodules.

3. **Spleen.** Enlargement of relatively moderate degree was common. (Average for series A., B. and D. 4.6 gm., average for controls 1.3 gm.) Tumour-like nodules were never encountered.

4. **Lymph nodes.** Apart from pale blue staining these structures did not shew morbid changes.

5. **Bone marrow.** This was uniformly pale slaty-grey to dull greyish-blue. Rarely it was faintly gelatinous.

**Intercurrent disease.** Sarcoptic infestation was encountered sporadically. This was treated with 'tetmosol' (Tetra-ethylthiureram monosulphide) with satisfactory results. Several animals were found to be lousy at necropsy. Rat bronchiectasis (Passey, Leese, & Knox, 1936) was a troublesome complication which forced the abandonment of the original series (not recorded here). In the present records this lesion also appeared but not to a serious degree, the controls shewed a rather lower incidence of it.

**Microscopical examination.**

**Liver.** There was a remarkable focal proliferation of cells in the portal tracts. In the initial stages this was fairly uniform throughout
These formations rapidly assumed the form of sharply demarcated rounded nodules which appeared as buds growing from an axial stem of portal tract tissue. Their origin appeared to be from pericytes in the adventitiae of the vascular radicles. The component cells conformed to a single type with the following characteristics. They measured 9-15μ diameter, rarely more, and were typically spheroidal or polygonal, but wedge-shaped or spindle-shaped when closely packed. The cytoplasm was non-granular, tending to be refractile, and oxyphile. Inclusions e.g. dye, haemosiderin, or debris, were common. The nuclei were trachychromatic and roughly oval. Detail was difficult to discern due to their density but they resembled shrunken twisted skeins. Mitoses were rare. Reticulin formation by these cells was the rule, at least in mature deposits. The formations usually increased with age but individual deposits outstripped others. Central cystic change and necrosis were common. In a small proportion of cases demarcation became poor, the growth rate increased and the deposits became tumour-like. The progression of the lesion was capricious in cases followed by biopsy in Series E. The cessation of dosage at 120 mgm. in seven of these animals was followed by advance to the tumour-like picture in two animals. In the remainder the appearance remained static or regressed.

Spleen. Enlargement was related only to general reactive hyperplastic change and engorgement or oedema. Haemosiderosis in the red pulp was seen occasionally. Tumour-like focal aggregates of macrophages were not encountered. (In all the spleens fields microscopically similar to Hodgkin's disease could be found; these were seen where septal connective tissue and megakaryocytes were in propinquity. The
resemblance is illusory, it did not mature at all).

Lymph nodes. In several cases the axillary nodes shewed partial replacement of lymphocytes by plasma cells and mast cells. Lesions comparable with the hepatic ones were never seen. An isolated finding in one node was diffuse fibrosis of the cortex.

Bone marrow. This was not consistently examined. Where sections were taken it was common to encounter scattered macrophages in the tissue, many contained dye.

Perivascular proliferation of cells like those in the liver lesions was constantly observed in the lungs, kidneys and omenta.

In all 12 attempts were made to transplant the tumour. Saline suspensions were made with aseptic precautions and injected intraperitoneally into adult male rats. Despite much care infections usually resulted, and eight of the animals died within three weeks. In the remaining four no trace of the inoculum was found at necropsy.

Series F. Mice receiving trypan blue.

Staining developed within a few days of the first injection and was exactly comparable to that observed in rats. On the other hand their condition remained normal. Animals were killed at 70, 100, 120, 170, 180, 200, 220, 250 and 300 (2 animals) days. The remainder survived in excellent health and were killed at 360 days. None were lost from the series and none died spontaneously.

At death faint general staining of the tissues was seen. The lymph nodes were readily found due to their faint blue colour, and were generally much easier to recover than the nodes in the rats. Ascites was not recorded and the livers shewed only a dark water-chocolate colour.
colour. Splenomegaly was absent.

Microscopical examination.

Liver. Circumscribed nodular hyperplastic aggregates of macrophage-like cells were observed in three animals (170 and 360(2) days). They were scanty and small in each case. In all animals after 70 days exposure to dye small haemopoietic foci were seen; most were localised in the sinuses, and a few were in portal tracts. In all the Kupffer cells were prominent due to dye content. Haemosiderin was very scanty in these cells.

Spleen. Haemosiderosis of moderate degree was common in the red pulp, but was not consistently related to the time of exposure.

Lymph nodes. Sinus catarrh was seen occasionally.

Kidneys. In all cases numerous dye granules were found in the first convoluted tubules of individual nephra. Focal interstitial or peri-vascular lesions were not encountered.

Lungs. Scanty perivascular and peribronchial deposits of small densely staining nuclei were seen. Dye and haemosiderin-laden macrophages were common in the alveoli.

Series C. Rats receiving vital new red.

Unlike trypan blue this dye did not appear toxic to rats in the dose used. All the animals remained outwardly healthy throughout the experiment, and none died spontaneously. Staining developed within three days of the first intra-peritoneal injection but partly on account of its colour it was difficult to detect unless controls were examined at the same time.

Six animals were killed after 27 fortnightly injections and six after 28 injections. One animal was accidentally killed in the first week of
the experiment, and two escaped, after almost a year.

At death very faint pink staining of the tissues was appreciable. The only conspicuous abnormality, which was seen in 7 animals, was enlargement of the juxta-caecal mesenteric lymph nodes. These were swollen and grey, and section revealed gelatinous material apparently occupying cysts in their substance. Ascites was not observed. Hepatomegaly and splenomegaly were absent, and the tâches laiteuses were not obvious. The average liver weight was 9.5 gm., controls 12.5 gm.; average spleen weight 0.9 gm., controls 1.3 gm.

**Microscopical examination.**

**Liver.** These were almost entirely normal. In two livers minute and scanty circumscribed aggregates of macrophage like cells were noted. The Kuppfer cells did not shew any increase.

**Spleen.** In four moderate haemosiderosis in the red pulp was noted, otherwise the glands were normal.

**Lymph nodes.** The peripheral nodes shewed sinus catarrh inconstantly. The changes in the caecal nodes were confirmed to be cystic change initiated in the lymphatic sinuses. The cellular conformation of these nodes remained normal.

**Lungs.** Rare aggregates of small macrophage-like cells were encountered in most.

**Kidneys.** Granules of dye were always seen in the tubule cells but rarely were they numerous.

**Comment on findings, and discussion.**

The most impressive finding is that when rats are injected with relatively small doses of the vital stain trypan blue at fortnightly intervals
intervals over a period of four to five months tumour-like nodules of distinctive structure develop in the liver in a considerable proportion of the animals (15 out of 23; 65%).

Progressive anaemia, emaciation, and weakness also result. Injection of the same individual doses with twice the frequency induces the same changes more rapidly.

In mice subjected to the same treatment these tumour-like changes in the liver are not produced to any extent, but anaemia may develop.

The growths produced appear to be constituted by a single cell-type which morphologically is a macrophage.

Rats given vital new red under the same conditions do not shew changes beyond inconstant haemosiderosis in the spleen.

In these experiments distinctive lymph node lesions were not observed in any of the animals and the 'Hodgkin-like' picture reported by previous workers was not reproduced. Occasionally slight alteration of lymph node cytology was encountered, plasma cells and mast-cells appearing to replace lymphocytes.

Interpretation of the hepatic lesion.

It is considered that it would be erroneous to designate the cellular growths obtained by chronic trypan blue poisoning as tumours, without qualification. Pathologists whose experience is chiefly derived from human morbid anatomy are well aware that to call a growth tumour is an award of considerable magnitude. In experimental cancer research, which is now virtually a separate subject, a less critical employment of the term neoplasm can be detected. In lower animals the sensitivity to neoplastic disease is nearly always much lower than in /man,
man, and constitutional disturbances e.g. illness, cachexia and anaemia are not so prominent. The anomalous brisk activity and good general health of 'cancer mice', which animals may have enormous tumours, is remarkable. These growths whilst legally cancer, are hardly the same in a moral sense. At microscopical level many experimental tumours present an indefinable appearance of artefact, at least to an observer familiar with human tumours.

In the present study the following features support the interpretation of the lesions as tumour. The growths present uniformity of the proliferating cell-type. There is comparatively little pleomorphism, certainly much less than is characteristic of a typical granuloma. This cell-type is atypical although it approximates closely in most respects to an histiocyte or macrophage. The possible affinities to erythroblast are mentioned but have not been specifically studied. In a proportion of cases more active proliferation is exhibited and simultaneously there is loss of demarcation of the growth which can be regarded as infiltration. There is in addition the phenomenon of progressive growth after removal of the presumed genetic/stimulus. Minor features supporting the view include the tendency to necrosis and degenerative change.

On the other hand most of the growths are sharply circumscribed and apparently indolent. Progression either with the continued exhibition of the dye or after its cessation is not invariable, some regression being common. The growths are confined to the liver and even where they appear malignant locally there is failure to metastasize. Finally attempts to transplant the tumours to other rats failed con-

/sistently.
sistent. This last phenomenon of serial transmission is characteristically a property of tumour, but the practical performance of the test is less certain than its theoretical possibility.

The property of producing reticulin neither reinforces nor detracts from the tumour hypothesis. The elaboration of reticulin by macrophages (histiocytes) is well authenticated, Curran, (1951); Nicol and Abou-Zikry, (1953).

In summary the growths are experimental tumours of characteristically low malignancy. They appear peculiar to the liver and are without close counterpart in human neoplastic disease. Morphologically they are sarcomata.

Assuming this view to be correct their genesis requires explanation. It might be held that repeated parenteral injection of a vital dye acting as a stimulant to the reticulo-endothelial system cells might result ultimately in neoplastic change. This has little to commend it. Generalised tumour production throughout the system was not found and the similar colloidal dye vital new red failed to cause similar growths. The total dye given was so small - generally under 0.5 gm., that in the latter case there was not even hyperplasia of the spleen. It is reasonable to infer that trypan blue may be carcinogenic in rats.

This dye is not generally credited with carcinogenic properties but this does not reflect final disposal of the possibility. The recent work of Simpson, (1952) and Marshall, (1953) following the Gillmans' studies, confirms that the dye evokes experimental tumours in rats. Boyland, (1952) has emphasized that in the present provisional list of chemical carcinogens powerful ones are chiefly considered, but that it is being learned that many less spectacularly potent agents also exist. The
effective dose of powerful carcinogens under optimum and standard conditions is very small. It has been demonstrated that as little as 0.2 mgm. of methylcholanthrene is capable of inducing skin papillomata in mice, Hieger (1953). Here trypan blue appeared effective in doses of about 120 mgm. In biological assay work disparities of this order viz. 1:600 are not exceptional, and this further credits the dye as a carcinogen under the present conditions of test. The restriction of the effect to one tissue site is a common phenomenon encountered in cancer research.

It is beyond the scope of the work to investigate the chemistry of this action. Some azo-dyes e.g. Butter Yellow, have been shewn to be carcinogenic and this property is ascribed to the presence of substituent methyl radicles in the amino-azo-benzene nucleus. o-tolidine, the basis of trypan blue contains two methyl radicles but is reported to be devoid of carcinogenic activity, Simpson (1952).

Two additional factors deserve mention. In general terms rats are readily susceptible to experimental sarcoma, Pullinger (1953), which has been induced by a variety of procedures. In contrast the mouse is refractory although it is typically susceptible to carcinoma.

This neoplastic propensity of rat connective tissue may even be evoked by culture and sub-culture of normal rat fibroblasts without the intervention of any recognised external factor. This almost incredible fact modifies ideas concerning the role of carcinogens in sarcomata of this animal and suggests that even the weakest carcinogens might be unexpectedly potent in this species.

Also it has long been known that the route of administration may determine
determine the action of a carcinogen. Superficial application is more likely to lead to carcinoma and parenteral injection to sarcoma.

Summary.

The prolonged administration of trypan blue (B.D.H.) in 10 mgm. doses at fortnightly or weekly intervals by the intra-peritoneal route induces a characteristic cellular proliferation in the hepatic portal tracts of rats. The type-cell of these lesions is a macrophage and the lesion is considered to be an experimental tumour.

The procedure also produces chronic poisoning and anaemia. Mice are refractory to this treatment and rats treated in the same method with vital new red do not shew similar changes.

In my hands the method has failed to produce lesions which simulate Hodgkin's disease.
SUMMARY AND CONCLUSIONS

In Part I an attempt was made to offer a brief and reasonably critical account of the inherent peculiarities of the structure and of the diseases of the reticulo-endothelial system. Particular emphasis was accorded to works which have served to integrate the concept of lymphoid tissue sarcoma. While it is doubtful whether knowledge is yet adequate enough to endorse this idea universally, it has the advantage of being a practical generalisation which simplifies the subject. The view that the morbid process, primary reticulosis, was covered by the generic lymphoid tissue sarcoma was also supported.

In Part II some account was given of the lymphatics and lymphoid tissue. In this outline attention was drawn to the mysterious and bewildering problems inseparable from the system. The structure of lymph nodes was given with observations on their development, involution, and possible neogenesis in adult life. From these studies it emerged that the full complement of lymph nodes in the locus examined was probably attained in adolescence or early adult life, and that fat replacement was the usual mode of atrophy. Attention was also drawn to the rarity of fibrosis in physiological nodes, except where it was the result of blood vascular hyaline change.

In Part III Hodgkin's disease was studied. In the introduction of this part of the work the historical aspect of the malady was recorded, with, it is hoped, advertisement of interesting and possibly less well-known facts about it. This was followed by a critical consideration on the nature of the disease and its morbid anatomy, the latter being illustrated
illustrated in part by analyses of the cases coming to necropsy at Glasgow Royal Infirmary over a period of fifty years. In this part also were the findings of a large series of biopsy specimens. Here endeavours were made to shew the microscopical variations in morphology in the lesion, and to demonstrate the affinities of other lymphoid tissue sarcomata with the disease. Within the resources available the generic lymphoid tissue sarcoma was established, and links between the better recognised variants were presented with a reasonable degree of conviction. In the necropsy series a detailed study of thirteen cases of Hodgkin's disease or reticulum cell sarcoma was offered. In these it was shewn that the favourite locus was lymphoid tissue, that complete systematisation was rare, and that metamorphosis to a more tumour-like lesion was common.

In Part IV two components of the Hodgkin's disease complex were studied in relation to general pathology. The view that fibrosis, an essential and inherent peculiarity of the Hodgkin's disease lesion, was represented in certain other morbid states was submitted. This was illustrated by brief accounts of some diseases where quasi-neoplastic features are shewn by connective tissue. Eosinophilia in tumours was also made the subject of investigation and revealed that the phenomenon, while possibly not so rare as might be expected, was not nearly so common as in Hodgkin's disease. Some evidence was found for the cyto-metaplastic origin of eosinophiles in Hodgkin's disease, but possibly due to the restriction to histological as opposed to cytological methods, the results were not highly conclusive.

In Part V an experimental attempt to reproduce the disease by
chronic trypan blue poisoning of rats and mice proved unsuccessful, although interesting results followed.

The main contention in this thesis has been that Hodgkin's disease is a neoplasm. Perhaps the following may influence the reader more convincingly than I have been able to do by so much work. The reasons for human beliefs depend chiefly upon Authority, Intuition, and Scientific Method. The last two have been exploited as far as I have been able; the foremost remains. As a junior student I saw a case of Hodgkin's disease first in the wards of the Late Professor Archibald Harrington, at Glasgow Royal Infirmary. I was chagrined at the doubt cast on its nature in the discussion which followed the demonstration; at twenty, one is very intolerant of obscure aetiology! On my return home I imprudently assailed my Father with the question at the dinner table, where even renal oedema was taboo. He was exceedingly angry. There was a dreadful silence, and then he relented - 'of course it is tumour, - but mind to whom you say that!' Nothing more was said. I submit that this terse pronouncement has been my most precious axiom, with deepest respect and affection.
This work was carried out during the tenure of a Dr. Foulis Memorial Scholarship in Pathology. The cost of the experimental studies and most of the photography and drawings was met by this grant. I thank the Trustees very sincerely for this award. It is my regret that the Trustees are anonymous because it is difficult to convey sincerity with conviction to an impersonal body, and to explain how much these funds have ameliorated the problem of surprisingly expensive research.

I am indebted to a great many people for gifts of material, for the loan of slides and for notes of cases. In particular I thank my Mother, Dr. W. Shaw Dunn, M.D., D.Sc. for the gift of my Father's collection of Hodgkin's disease slides which provided the first nucleus of this work.

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My own colleagues in the Department of Pathology at Glasgow Royal Infirmary, who must have come to regard this protracted work as the veritable Reticulum of Penelope, have been unstintingly generous in shewing me material and drawing my attention to cases. It is a very sincere pleasure to thank Dr. Alice Marshall, Dr. Alastair Currie, Dr. Robert Curran, Dr. Robert Patrick, Dr. William Crane, Dr. Bruce Woodger and Dr. Peter Pullar.
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The staff of the Library of the University of Glasgow have given me most kindly assistance, and have on many occasions obtained for me rare or comparatively inaccessible texts.

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Finally I thank my wife, Dr. Maida Shaw Dunn, for her incredible patience throughout the prolonged period of this study and the small members of the family for whom 'Daddy's Hodgkin' was virtually Pandora's box.
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20. Scyphomantic
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23. Sternberg: pseudo-leukemia

41. MacCallum: death But no Swine
38. Holoma
49. Identical for unidentical
68. lymph: frog - Klein Barnes
97. Immolation
227. atavistic

266. time cosmopolis
### VOLUME II

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APPENDIX A.

BIOPSY SERIES PROTOCOLS

The pages are numbered to correspond with the serial number of each biopsy; where the records exceed one page the appropriate serial numbers are retained and followed by the letter 'a'.

Illustrations.

Volume III Figs. 45-116.

Volume IV p.20-57.
Established Hodgkin's disease in which all diagnostic criteria are met. There is appreciable persistence of normal structure in one of the nodes, it is represented by sub-capsular lymphoid tissue and a few survival nodules in the deeper node substance. The capsules are irregularly thickened, especially at the bases of the trabeculae. Concentric re-formation and invasion are absent. Fibrosis is moderate, perivascular lamellation and coralline deposits are the chief expression. The cellular moiety of the new tissue is classical; the proliferating reticulum cells are seen in syncytial form in places, multinucleate giant-cells are numerous, often 2 per H.P. field. Eosinophilia is pronounced. Mitoses average 1 per H.P. field. Necrosis is confined to scattered individual cells. There is faint lobulation of the nodes and the vascularity is definitely low.

Diagnosis. Hodgkin's disease, Typical.
2. Serial 2.


There is pronounced fibrosis of the capsule and one large trabecula, small coralline deposits are seen below the capsule. The cellular tissue is curiously uniform, most of the cells are very slightly immature lymphocytes and they are distinctly sparser than normal. A diffuse increase of reticulum cells in present, they are uniformly scattered. Plasma cells are seen but eosinophiles and multinucleate giant-cells are absent. The fibrosis and general simplification are in keeping with Hodgkin's disease but the lack of pleiomorphism etc. are against it. Rate of growth is very slow.

Diagnosis. Indeterminate ? benign Hodgkin's disease. (Further sections might have elucidated this).

Established Hodgkin's disease in which all diagnostic criteria are met. The lesion here shews a peculiar but remarkable characteristic, the exodic growth phenomenon. Centrally in the node there is a uniform field of typically pleiomorphic Hodgkin's tissue but this does not quite reach the capsule, a thin rind of small lymphocytes with a few physiological reticulum cells being interposed. This attenuated lymphoid tissue is being driven against the inner aspect of the capsule. A few survival nodes of lymphoid tissue are also notable deeper in the new tissue. The capsule is poorly preserved (artefact) but does not appear much thickened. Trabeculae are not present. Fibrosis is diffuse - of synplasmic transformation type; perivascular and coralline fibrosis are slight. Fibrils and fibrillae constitute most of the collagen. Reticulum cells are conspicuously increased and are chiefly of the morbid mononuclear variety, some shew gradations to fibroblast resulting in a subtle medley of both cell types. Typical multinucleate giant-cells are scanty but Dorothy Reed mirror-image type are common, lobulated nuclei are also frequent. Eosinophiles are numerous and mostly of blood type. Plasma cells and neutrophils are identified. Mitoses are few and occasional areas of necrosis are noted.

The second specimen is slightly more mature and the fibrosis is more evident; however, even here the entire node is not involved. Note. A few fat cells are present in this node. This is a rare finding.

Diagnosis. Hodgkin's disease, Typical.
Early established Hodgkin's disease in which most diagnostic criteria are met. Under very low power a faint yet definite coarse segmentation is discernible, the exodic growth phenomenon is apparent as well. Survival follicles and nodules are present. The capsule where present is thickened grossly, true trabeculae are absent. Fibrosis is well established, it is seen in the form of a vaguely whorled mesh in the segments described. Coralline deposits and perivascular lamellation are also present. Mononuclear reticulum cells are relatively scanty, multinucleate cells appearing more common. Distribution of the latter is uniform. Eosinophiles and other granulocytes are rare. Mitoses are few, necrosis is only represented on an individual cell basis.

Diagnosis. Atypical Hodgkin's disease. Eosinophiles are distinctly few and pleiomorphism is less than usual. It might conceivably be confused with lymphosarcoma but not after close scrutiny.

This is a reticulum cell sarcoma of fairly uniform cell type—close to Gall and Mallory's clasmaticytic stem-cell lymphoma. The cell differentiation direction is wavering in places with bias towards lymphoid cell type. The capsule is not thickened but the concentric re-formation phenomenon is well shewn. Normal lymphoid tissue is totally absent. Fibrosis is present, it is strongly reminiscent of Hodgkin's disease fibrosis too. Diffuse fibrillary and fibril deposits, occasionally assuming the coralline form are notable. The vessels inconstantly shew lamellation. Typical multinucleate giant cells are absent but small forms occur. Eosinophiles are absent. There are areas of necrosis with local polymorph response. Mitoses are very numerous, far beyond the rate seen in Hodgkin's disease.

Diagnosis. Reticulum cell sarcoma, sclerosing in similar fashion to Hodgkin's disease.

Classical Hodgkin's disease, fairly early. The new tissue is disposed in vague segmental fashion in places and occasional survival aggregates of lymphocytes are present in the interstices.

The capsule is very incomplete due to artefact but thickening is seen locally and also infiltration with the new tissue. Connective tissue swathes of crescentic outline closely simulate trabeculae but the loss of capsule is too extensive to substantiate this. Fibrosis is marked, it is chiefly fibrillary and of the synplasmic transformation variety. Perivascular lamellation is present in addition.

The new tissue is hypocellular due to marked loss of lymphocytes. Reticulum cells are fairly numerous, many being indeterminate reticulum cell/fibroblast types. The giant cells are classical. Eosinophiles are very numerous indeed. The remaining characteristic cells are represented. Mitoses average about 1 per H.P. field and necrosis is absent.

Diagnosis. Hodgkin's disease, Typical.
Forsyth. Ward XX. Lymph nodes. History unknown. The first sections shew very good typical Hodgkin's disease, the second series from another node present an uncommon variant.

In the former all vestige of normal lymphoid tissue except an occasional aggregate of small lymphocytes has gone. The capsule is not preserved unfortunately, but new trabeculae enclosing the Hodgkin's disease segments are discernible, this change recalls the results in sclerosing Brill's disease. Fibrosis is pronounced and all varieties are exhibited in some degree. The reticulum cell proliferation is well shewn, the cells are chiefly of the morbid mononuclear type. The giant cells are typical. Eosinophiles are extraordinarily numerous; in addition small mononuclear cells with lymphocyte nuclei and eosinophile ground-glass cytoplasm are seen. In many cases actual eosinophile granules are present in these and a lymphocytic origin of some of the eosinophiles seems very likely. Plasma cells are common but polymorphs are rare. Mitoses are numerically above average but there is no necrosis. In the second set multinucleate giant cells of Langhan's type are the commonest giant cell. Up to 60 nuclei were counted in one cell. There is not any evidence of tuberculosis, e.g. follicles, caseation etc.

An interesting feature is seen in these specimens regarding the relation of reticulin to collagen. The correlation of the silver impregnations with the routine sections is poor, they appear quite unrelated.

Diagnosis. 1. Hodgkin's disease, Typical.
McIntyre. C.C.C. History unknown.

The specimen is a lymph node shewing late Hodgkin's disease.

There is not any persistence of normal structure and the capsule is incomplete.

Fibrosis dominates the picture; centrally there is a mass of almost avascular and acellular coralline collagen which peripherally merges with a mantle of active Hodgkin's tissue, the latter is imperfect being traversed by trabeculae. The new tissue is characteristic except for the paucity of eosinophiles but this is common in mature lesions. Mitoses are very few. The appearances support the idea that the lesion starts centrally and drives out. The lesion is subtle in the preparation but the column behind the furled colours is a unit of the same army.

Diagnosis. Hodgkin's disease, Typical late.

In one of the nodes, the smallest, there is some surviving lymphoid tissue; follicles and sinus are represented. This persistent tissue is limited to the extreme periphery of the node. The node capsules exhibit irregular thickening and hyalinisation. Concentric re-formation and invasion are absent. Trabeculae are poorly developed and consist of blunt fibrotic wedges. Fibrosis is present but fairly early, it is chiefly in fibrils and fibrillae related to the mesh. Perivascular deposits and coralline formation are little in evidence. The reticulum cell increase is characteristic and there is copious production of morbid mononuclear types. Typical Hodgkin giant-cells are present in about the usual number and there is vague topical aggregation of them locally. In one section a typical Langhan's giant-cell is seen apparently encircling an Hodgkin's giant-cell. This is extremely interesting because it suggests that the cells are quite distinct—a mutual antipathy existing between them. The Langhan's cell can reasonably claim to be altruistic in function and although one swallow does not make a summer it is a tiny evidence in favour of the neoplastic character of the Hodgkin cell. (This remarkable occurrence has not been seen again). Eosinophilia is typical and the remaining characteristic cells are present. Here the polymorphs are definitely related to focal necroses. Cellular activity is of the normal order.

The exodic growth phenomenon is very well shewn.

Diagnosis: Hodgkin's disease, Typical.

Seven lymph nodes are available for study, and several striking findings are notable. In one node fully established typical Hodgkin's disease is present and yet perfectly normal physiologically active lymph node tissue containing follicles with pale centres and sinuses is still present in the same gland. Under very low power a vague frontier between the two tissues is just visible giving an impression that the new tissue is driving into and overwhelming the old. Under ordinary low power demarcation becomes guesswork. The exodic growth phenomenon is seldom so emphatically demonstrated and a further fact emerges - the only reaction of the normal tissue is a general reactive hyperplasia, not a 'lymphoid hyperplasia' which is usually postulated as the incipient change. In five other nodes the typical Hodgkin's disease picture is completely characteristic, all criteria are fully satisfied. In the remaining node the multinucleate giant cells are unusually numerous giving the appearance of Warthin's Hodgkin's sarcoma but eosinophiles are still very copious. This variation in picture is seldom able to be confirmed in specimens from the living since diagnostic resections are limited in extent. The phenomenon is however amply demonstrated in full necropsy studies.

Diagnosis. Hodgkin's disease, Typical.
McFarlane. Ward XX. Two lymph nodes, incomplete. History unknown.

There is not any persistence of normal lymphoid tissue in either specimen. The capsules show pronounced fibrous thickening and hyalinisation. The fibrosis is advanced and obscures evidence of concentric re-formation, the trabeculae are particularly massive. A coarse net is formed which segments the node. Perivascular lamellation is the most obvious display of collagen formation, the other forms including coralline deposits are difficult to recognise. The reticulum cell proliferation is chiefly productive of morbid mononuclear types and Hodgkin's giant-cells are few. Eosinophilia tends to be focal. It is pertinent that these cells are mainly located in areas where lymphocytes persist. Plasma cells are more numerous than usual and transitions between these and lymphocytes are common. Judged by the number of mitoses growth is slow. The growth in irregular large islets is still retained despite the maturity of the lesion.

Diagnosis. Hodgkin's disease, Typical, late.

These specimens are poorly preserved, attempts to restain one of the slides met with indifferent results. Where the capsules are preserved there is uniform thickening and hyalinisation. The trabeculae are thickened and in their immediate vicinity the appearances of typical Hodgkin's disease are seen. Beyond these limits the nodes are replaced by a uniform small round cell tumour which corresponds to mixed reticulum cell/lymphosarcoma type. There is total absence of eosinophiles in the homogeneous tumour portion and mitoses are numerous. The exodic growth phenomenon is particularly well illustrated in one node, the peripheral mantle of crushed lymphoid tissue containing germinal follicles which have been flattened by pressure.

Diagnosis. Mixed reticulum cell and lymphocyte series sarcoma, corresponding to intermediate cell type of Ehrlich and Gerber (1935); with fields adjacent to the trabeculae morphologically identical with Hodgkin's disease.

History unknown.

In each specimen there are fairly numerous survival follicles of lymphoid tissue, most contain normal active pale centres. Their situation is not limited to the periphery of the nodes. Capsular thickening is apparent in only one gland. The trabeculae are very poorly developed. Fibrosis is extensive, perivascular lamellation is seen but most of the collagen is present as a diffuse fibril and fibrillar mesh with scar-like areas of unusual intensity. Reticulum cells are numerous, morbid mononuclear examples are about average in number but the majority resemble epithelioid types. Typical Hodgkin giant-cells are present but there are more which correspond closely to Langhan's type. A form with a complete ring of peripheral nuclei is common. Eosinophiles are present in adequate numbers but neutrophiles are few. The lesion appears curiously inactive. Mitoses are hard to find.

Diagnosis. Atypical Hodgkin's disease, Early but established. Abnormal multinucleate giant-cells (Langhan's type). Bears a faint resemblance to endothelial tuberculosis. Origin in the loose lymphatic tissue probable.

This is a very interesting set. In several of the nodes there is a rind of persisting normal lymphoid tissue including germinal follicles with pale centres and fragments of marginal sinus. The capsules of all specimens are thickened. In one mode the growth pressure of the new tissue has caused massive infarction. Nearly all the Hodgkin's tissue is dead, except for a thin sub-capsular rind in which some normal (reactive) lymphoid tissue is identified. In the largest node new trabeculae are forming giving a coarse follicular pattern superficially like Brill's disease but the segmentation is less regular. Fibrosis is chiefly seen as perivascular lamellae and capsular thickening but incipient diffuse fibril formation is in progress. The reticulum cells are unusually numerous and although most are of morbid mononuclear type small 2–3 nucelated giant-cells are common. Typical Hodgkin giant-cells are present. Eosinophilia is diffuse and heavy, other granulocytes are rare. Plasma cells are very scanty. Mitoses are about average.

Diagnosis. Hodgkin's disease, rather more cellular than usual. Some affinity to Brill's disease but regard as typical.
Allan. XXXIV. Single section of lymph node. History unknown.

There is not any remaining normal tissue. The capsule is grossly thickened and hyalinised. Coarse segmentation by pseudo-trabeculae is pronounced. Fibrosis is chiefly by perivascular accretion and is gross. Coralline deposits are particularly marked. The reticulum cell proliferation is unusually florid, most approach the macrophage type as regards nucleus but are syncytial. Typical Hodgkin giant-cells are present but the majority are intermediate between this variety and Langhan's type. They are grotesquely angular and irregular, recalling the foreign body giant cell. Eosinophiles and plasma cells are very scanty. Neutrophiles are absent. Mitoses are rare but occasional areas of necrosis are notable.


History unknown.

Rare survival follicles and islets persist. Elsewhere there is a very wide replacement of the node tissue by a rather homogeneous cellular Hodgkin's tissue. Capsular thickening and hyaline change are present and some concentric re-formation is detected. The new tissue despite its uniform appearance low power meets all criteria for its recognition, eosinophilia is unusually marked. The fibrosis is patchy and mostly of the coralline deposit variety. Mitoses are absent.

Diagnosis. Hodgkin's disease, accept as typical but rather uniform and cellular.
Lee. VI. February 1913. A single lymph node. History unknown.

A few fragments of the marginal sinus tissue persist, the littoral cells are laden with haemosiderin. The capsule is grossly thickened and hyaline change is advanced. Concentric re-formation is notable, several crescentic swathes of new fibrous tissue have formed. The few trabeculae present are thickened by fibrotic accretion. Perivascular lamellation is pronounced but fibrosis in the lymphoid tissue is limited to small scar-like areas of coralline type. The reticulum cell proliferation is about usual, but large lymphocytes of immature variety still predominate. Typical Hodgkin giant-cells are present but the mirror image type is much commoner. Eosinophilia is mild, necrosis and mitotic figures are scanty.

Diagnosis. Hodgkin's disease, typical but rather less pleiomorphic than usual.
Inglis. XXXIV. Dated February 1913. Single lymph node. History unknown.

This is late Hodgkin's disease. The fibrosis is overwhelming. Definition of the capsule and trabeculae is no longer possible and the new tissue itself is seen in survival islets in interstices of the collagen. In these areas it is typical and easily recognised. The reticulum cells are chiefly of the morbid mononuclear type but Hodgkin giant-cells are present too. Eosinophilia persists and a few neutrophiles are seen also.

Diagnosis. Hodgkin's disease, Typical, late.
Callaghan. XX. Dated 13th June 1913. Three lymph nodes.

History unknown.

This is primarily a mixed lympho-reticulo sarcoma (intermediate type of Ehrlich & Gerber (1935)). There is not any survival of normal lymphoid tissue. The capsules are thickened but also infiltrated and over-run. Concentric re-formation is present. The tumour is growing in coarse ill defined nodules 1-2 mm. diameter which are defined by swathes of connective tissue. Fibrosis is also notable in among the tumour cells giving the descriptive 'alveolar sarcoma' picture. Hodgkin's giant-cells are present but very sparse. Eosinophilia is absent and plasma cells are not seen. Mitoses are numerous, 2-3 per H.P. field; necrosis is confined to individual cells.

Diagnosis. Intermediate type reticulum cell sarcoma, Fibrifying. Some features of Hodgkin's disease, occasional fields would pass for it.

This is nearest to follicular lymphoma but fibrosis is taking place in it. The false follicles are irregular in size and shape. Their pale centres present an unusually pleiomorphic cell picture. Many fields indeed would pass for Hodgkin's disease. The extra-follicular tissue is even closer to the latter in morphology. Fibrosis is present round many follicles, sometimes in their centre, and perivascular lamellation is marked. Most of the proliferating reticulum cells are of the morbid mononuclear type but scanty typical Hodgkin's giant-cells are visible as well. Eosinophiles, plasma cells and neutrophiles are very sparse. Mitotic activity is well above Hodgkin level.

Diagnosis. Brill's disease, sclerosing; very close to Hodgkin's disease.
Identity unknown beyond M.R.I. 27:243. History unknown. Several sections of four different lymph nodes. The appearances are the same in each.

This is an interesting example of a rare variant of Hodgkin's disease. Normal lymphoid tissue is absent. The capsules show slight but definite fibrous thickening of hyaline character. The exodic growth phenomenon is displayed well and in a few places the new tissue is being driven into the substance of the capsule. Fibrosis is chiefly expressed in the formation of extensive fine fibrils, with occasional spidery scars. Perivascular deposit is seen round some vessels but this localisation is inconstant. Coralline deposits of collagen are absent. The reticulum cell proliferation is meagre, morbid mononuclear types are identified but are few. Multinucleate giant cells are very numerous and the great majority are of Langhan's type, the typical Hodgkin cells being hard to find. Eosinophilia is pronounced, but other granulocytes and plasma cells are few. Mitoses are rare and necrosis is absent. The commonest cell present is the small lymphocyte and it appears normally mature.

Diagnosis. Atypical Hodgkin's disease, Langhan's type multinucleate giant cells. Some reduction of pleiomorphism.
Identity. 30:200. (Path. Lab; University of Glasgow Label.) History unknown, believed to be from a patient in Manchester Royal Infirmary. A large slide presenting sections of five apparently different lymph nodes.

This is of particular interest in two respects. Normal lymphoid tissue is identified in nodes shewing established Hodgkin's disease, and metamorphosis to Hodgkin's sarcoma (Warthin's reticulo-endothelioblastoma) is seen in three of the nodes.

One node, V, which is adjacent to IV, shews only reactive change with much macrophage production. There is haemosiderin storage in some of these cells. Node III shews late Hodgkin's disease, the only irregularity being a low eosinophil content. Node I shews Hodgkin's disease and Hodgkin's sarcoma, between which transition is clearly discernible. Nodes II and IV shew almost pure Hodgkin's sarcoma which is becoming uniform. In this case the sarcomatous transformation is producing a tumour of unquestionably malignant character which morphologically is a reticulum cell sarcoma of the variety clasmatocytic stem cell lymphoma of Gall and Mallory. The eosinophils have almost disappeared from these lesions. The growth is diffuse in the pulp and the sinus tissue appears remarkably intact in some places. (This case is more fully described under Hodgkin's sarcoma q.v.)

Diagnosis. Hodgkin's disease, Typical. Hodgkin's sarcoma. (Reticulo-endothelioblastoma of Warthin), Transition between the two appearances is convincing.
A single lymph node 1.5 x 1 cm., from the axilla.

This is an interesting variety of atypical Hodgkin's disease which possibly forms a link between the benign Hodgkin's disease of Harrison and Hodgkin's disease; it is undoubtedly nearer to the true disease. There is persistence of normal (reactive) lymphoid tissue at the subcapsular periphery and a distinct exodic growth phenomenon is discernible. The capsule is irregularly thickened and attenuated, it is not invaded. Trabecular lobulation is not apparent. Fibrosis is intense at one pole of the node and is mostly of coralline variety though perivascular reticulum cells are present and a few typical Hodgkin giant cells are seen. Eosinophils are sparse and so are plasma cells and neutrophils. Mitotic figures are rare and there is not any necrosis. Normal mature small lymphocytes are very copious except in the fibrosed area.

Mrs. Margaret Wright aet 59. Ref. G.R.I. 1336:33. 3rd October 1933.
Following trauma two years ago patient noticed masses on her head.
14 months ago the post-auricular chain of lymph nodes began to enlarge, and eventually others at the base of the neck enlarged as well.

Cervical lymph node.

This is late typical Hodgkin's disease. The lesion is particularly fibrous and it is not possible to identify the capsule with certainty. The fibrosis is mainly in compact swathes of fibrils which segment the node in irregular fashion. Perivascular lamellation is also seen. The cellular areas of the lesion are located in these collagenous casemates. The reticulum cell increase is slight, but morbid mononuclear and Hodgkin giant cells are identified. Eosinophilia persists but is restricted to the more cellular portions. Plasma cells and neutrophils are scanty. Mitotic figures are very rare, necrosis is shown only by occasional individual cells.

Diagnosis. Hodgkin's disease, Typical, late.
Inguinal lymph node.

This is an example of reticulum cell sarcoma which exhibits certain features characteristic of Hodgkin's disease.

The bulk of the node is occupied by tumour, at one pole normal reactive lymphoid tissue is seen being driven against the inner aspect of the capsule. On the lymphoid tissue side of the line of contact of tumour and normal tissue, a crescent of fibrous tissue is forming. The node capsule is thickened and hyaline and, significantly, it is not invaded. The tumour is a stem cell lymphoma type of which the nuclei shew a definite notching of their membranes which are pachychromatic. Aggregates of tumour cell nuclei which simulate Hodgkin giant cells are common but no cytoplasmic component is present. Mitotic activity is high, 3 per H.P. field. Fibrosis exists in the growth both as a coarse mesh and a fine filigree; and in some areas coralline deposits are present. A reticulum impregnation supports the interpretation of sarcoma and not that of anaplastic carcinoma. High power scrutiny reveals many fields which would pass for Hodgkin's disease.

Diagnosis. Reticulum cell sarcoma. Shews affinities to Hodgkin's disease viz;— failure to invade the capsule, Hodgkin type of fibrosis, focal Hodgkin morphology.
It is difficult to decide what is the most appropriate category for this lesion. Morphologically it is a fibrifying reticulum cell sarcoma in which an unusually perfect follicular pattern is present. The cellular deposits are mostly spheroidal or deformed spheroidal and correspond very closely with the false follicles of follicular lymphoma. These are set in a matrix of dense collagenous fibrous tissue which is made up of compacted coralline variety elements. In some areas mixtures of cellular and fibrous parts exist which are almost indistinguishable from Hodgkin's disease, even to the presence of Hodgkin giant cells. The growth is not slow growing, mitoses are common; but at the same time the casemates of collagen seem impenetrable. Massive necrosis presumably due to infarction is seen at one part. While perifollicular fibrosis is common in late follicular lymphoma which remains relatively benign, the type of fibrosis seen here is unlike it. Further when such a lesion becomes more malignant the fibrosis is scanty as a rule and the follicular pattern fades.

Diagnosis. Reticulum cell sarcoma, presenting an unusual pattern of fibrosis. Related morphologically to Brill's disease and Hodgkin's disease.

This is an example of Brill's disease in a lymph node. The lesion is of considerable standing and fibrosis is pronounced. The capsule is not preserved or no longer distinguishable. Fibrous shells have formed round the giant follicles and are as much as 100 micra thick in places. Perivascular lamellation fibrosis is marked and scattered coralline deposits are frequent. The tumour is apparently of slow growth, mitoses are not frequent. Deep in the node towards one pole an infarct of some considerable age is present. It measures 1 cm. diameter, (specimen 3 x 1.5 cm.)


This is an example of Brill's disease in which the follicles are small and the follicle cells more lymphoid than reticular in their affinities. The lesion must be regarded as malignant; there are transgression of the capsule, deposits outwith the capsular breaches which reproduce the tumour pattern, and infiltration of the fat. Definite, but unimpressive, fibrosis is seen round some follicles. This is not apparently related to the node reticulum being compressed but to small blood vessels which are undergoing hyalinisation. Further fibrosis is seen round larger blood vessels and also in the pulp (follicles).

Diagnosis. Brill's disease, undergoing malignant change to small cell lymphosarcoma. Tendency to sclerose slight, but present.
Ref. Ayr County Hospital. 3:37. Holograph (J.S.D.) "? plump cellular stage" on label. A small lymph node (11 x 3 mm). Site unknown.

Small lymph nodes are nearly always inadequate for reliable diagnosis of Hodgkin's disease. This example is more informative than most but the assessment is provisional.

There are areas of normal tissue remaining, these consist of small deposits of loose lymphatic tissue round the hilar intrusion and portions of the medullary lymph sinuses. Elsewhere the pulp and follicles are replaced except for tiny survival islets of lymphocytes. The capsule is definitely thickened and there is a perceptible increase of connective tissue round arterioles in the node substance. The new tissue is composed of a uniform crowd of slightly abnormal reticulum cell nuclei whose cytoplasmic component is not discernible. The nuclei are larger than normal, oval, and tending to become vesicular; oxyphilic nucleoli are common. They approach the morbid mononuclear cell of Hodgkin's disease, but most are not large enough to qualify for this characterization, a very few may be considered typical, but Hodgkin giant cells are not seen at all. Eosinophils are present but sparse. Plasma cells and neutrophils are very scanty. The reticulum cells present many mitoses.

Cannon. Ref. Greenock. 10/37. A lymph node from the posterior triangle of the neck of a boy aged twelve years.

The node is a small one (10 x 4 mm.) and has been sectioned at four levels.

At one level the appearances are within normal considering the patient's age. A further level shews only one morbid area and the remaining two are diffusely affected.

The capsule is of normal thickness. The exodic growth phenomenon is evident and the normal architecture of the node is disappearing. In its place ill-defined lobules approx. 1 mm. diameter are appearing. These are composed predominantly of normal lymphocytes but scattered morbid mononuclear reticulum cells and typical Hodgkin giant cells are present. Incipient fibrosis is present; it is seen mainly alongside the trabeculae and as diffuse fibrillae. Eosinophils are absent and so are plasma cells and neutrophils.

Diagnosis. Hodgkin's disease, early. Pre-eosinophilia stage.

Note. This decision is arrived at after many examinations at intervals over several years; it is admittedly partly intuitive. The tissue has the capriciously disordered look of Hodgkin's disease.

The specimen consists of two small nodes with adjacent areolar tissue and an incomplete larger node. The intact nodes show general retention of their architecture with unusually marked sinus hyperplasia of the solid type. The pulp shows the enhanced heterogeneity of cells characteristic of reactive change but occasionally large tumour-like nuclei of reticulum cell type are appearing. The damaged node shows a picture which may be regarded as extremely cellular Hodgkin's disease or reticulum cell sarcoma. Here the capsule is of normal thickness where it is preserved, but it is being over-run by the tumour. Fibrosis is present, all varieties except coralline deposits are seen, but the amount is very small. The reticulum cell proliferation is extraordinarily profuse and many of the cells are typical morbid mononuclears. In addition Hodgkin giant cells are present in large numbers; some giant cells conform closely to Langhan's cells in the spatial arrangement of their nuclei but the resemblance is otherwise superficial. Eosinophils are present, but rather below the concentration typical of Hodgkin's disease. Plasma cells are present and neutrophils are notable in focal areas. Mitoses, including spectacularly abnormal ones are common, up to 7 per H.P. field. Necrosis is absent.

Diagnosis. Atypical Hodgkin's disease, very cellular, tumour-like, or reticulum cell sarcoma. (The retention of eosinophilia is anomalous).

This is an example of either slightly atypical highly cellular Hodgkin's disease or reticulum cell sarcoma. The capsule is slightly damaged but is thickened slightly elsewhere. It is over-run at several points by the growth. The new tissue is arranged in ill-defined follicles which are unequal in size and often mutually distorted. Fibrosis is slight, it is visible round blood vessels and as tenuous deposits of fibrillary and fibril type. The predominant cell type is a morbid mononuclear reticulum cell among which many typical Hodgkin giant cells are scattered. Eosinophils are distinctly numerous and plasma cells are also common. Neutrophils in small aggregates and scattered diffusely are frequent. The mitotic count is very high, as many as 20 being visible in some H.P. fields. Necrosis is limited to occasional individual cells.

It is really equivocal whether this is more deserving of one or other of the alternative designations suggested. If microscopical morbid anatomy means anything this lesion is among the most convincing examples of the essential identity of the two processes.

Diagnosis. Atypical Hodgkin's disease, Cellular; or reticulum cell sarcoma.
This is an example of atypical Hodgkin's disease in which eosinophilia is lacking and in which the usual capricious pleomorphism is obscured by the persistence of large numbers of small lymphocyte series cells. The capsule is imperfectly preserved due to artefact, occasional relics of the marginal sinus are recognisable. Concentric re-formation of this envelope is well shown at one pole. The new tissue is irregularly follicular in arrangement with a distinct pseudo-lobulation. Fibrosis is well established, perivascular lamellation, fibrillary and coralline varieties are all seen. Morbid mononuclear reticulum cells are present but not very numerous. Most of the reticulum cell nuclei are normal in appearance. Hodgkin giant cells are present and several shew central aggregates of their nuclei, recalling the Touton giant cell. Eosinophils are not seen. Plasma cells and neutrophils are scanty. Mitotic figures are rare. Almost mature small lymphocytes are very numerous and obscure the pathognomonic picture in places.


This is early Hodgkin's disease in which the new tissue is seen only in focal areas of the pulp of the node. Much of the anatomical cortical tissue is normal and follicles with pale centres are numerous. The cells of the last locality are not visibly contributing to the new tissue at all.

Fibrosis is already present in all forms except coralline deposits. Morbid mononuclear reticulum cells are conspicuous and there are fairly numerous Hodgkin giant cells. Eosinophilia is marked, but plasma cells and neutrophils are very few. Mitosis is seen but is infrequent.

The specimen is an incomplete enlarged lymph node (27 x 15 mm.) It is an example of pleiomorphic reticulum cell sarcoma with considerable affinity to Hodgkin's disease. The whole tissue is morbid. The capsule is grossly thickened, up to 1 mm. in places and abstracted deposits of the tumour are embedded in it. Several densely fibrous trabeculae of great width are present. Fibrosis is chiefly expressed in the above forms but some perivascular deposit is also present. The tumour is composed of large morbid reticulum cells with discrete cytoplasmic bodies. Synplasmic growth is visible in small areas. The nuclei are trachychromatic, assume bizarre forms, and frequently exhibit oxyphile nucleoli. Hodgkin giant cells are identified, they are rather below the usual size. Eosinophilia is absent. Mitoses are not numerous - many apparent figures prove to be pyknotic fragments. Massive necrosis is present in several areas, at one part forming a crescent 6 x 2 mm. Deposits of tumour are present in lymphatic and blood vessels.

Diagnosis. Reticulum cell sarcoma, pleiomorphic. Affinities to Hodgkin's disease (fibrosis, cell forms).

This is a follicular lymphoma in which the false follicles are small and composed of small deeply staining cells, recalling the solid follicles of normal lymphoid tissue. The capsule is imperfectly preserved but appears to be infiltrated. Fibrosis is present; the component round arterioles is surprisingly large, and coralline deposits are notable in a few areas, apparently being laid down in the follicles. Reticulum cell proliferation is evident, morbid mononuclear types are identified and a very few Hodgkin giant cells are visible. Eosinophilia is absent. The mitotic activity is about 2 per H.P. field.

This is a lymph node which has been partly atrophic before the lesion developed. There is a formation of large follicles at the periphery of the convex border but elsewhere a physiological sinus hyperplasia is the chief feature.

Normal physiologically active lymphoid tissue is present in the concave side of the node. The sinuses are tightly packed with normal littoral cells many of which contain erythrocytes. The capsule is thickened but chiefly on the convexity of the node, which is a normal finding in involuting nodes. There is moderate fibrosis, this is seen in the trabeculae and as coralline deposits. Almost all the latter can be accounted for by vascular fibrosis, i.e. it is not integral to the lesion. The large follicles are not uniform in size or outline, their pale centres are often composite nodules of cells including reticulum cells, lymphocytes and haemosiderin-laden macrophages. Their cellular heterogeneity is in contrast to the pale centres of the false follicles in Brill's disease. Small Hodgkin giant cells are present in very small numbers and an occasional eosinophil is noted. Plasma cells are common. The mitotic activity of the reticulum cells is low, below 1 per H.P. field.

Diagnosis. Indeterminate. ?? Early (focal in node) Brill's disease. Regarded as probably reactive.
38. Serial 38.


This is an example of follicular lymphoma which is growing as a large circumscribed nodule in the lymph node and shews pronounced topical eosinophilia and perivascular fibrosis.

A crescentic mantle of crushed normal (reactive) lymphoid tissue can be made out round most of the tumour nodule. The capsule is normal.

In the tumour nodule many false follicles of varying size and shape have developed. Most are of the solid variety the predominant cell being the small lymphocyte. Morbid reticulum cells are present but are few. A very few Hodgkin giant cells are seen. Eosinophils are distinctly numerous, comparable with an average case of established Hodgkin's disease. Slight fibrosis is seen, it is mainly perivascular with a small fibrillar moiety. Mitotic activity is moderately high, up to 4 mitoses per H.P. field.

This is lymphadenoid tissue probably lymph node almost entirely replaced by reticulum cell sarcoma.

At one point of the periphery nerve tissue is over-run by the tumour. There is fibrosis which segments the growth into alveolar sarcoma, in addition perivascular and fine fibrillary components are seen. The tumour corresponds to the clasmatocyte stem cell lymphoma of Gall and Mallory, occasional Hodgkin giant cells are seen but their cytoplasm is small in amount. Eosinophils are practically absent. Plasma cells and cells of acute inflammation are common. Mitoses are very numerous.

This is Hodgkin's disease of highly cellular uniform type, it is distinctly tumour-like in appearance.

The capsule shows local thickenings, chiefly at the bases of trabeculae, elsewhere it is thin. The exoctic growth phenomenon is well shewn but the peripheral mantle of lymphoid tissue is in process of being taken up in the tumour process. Fibrosis is present but in an unusual form. A delicate fibril mesh of open type has formed and is joined by very small dibrillae from the tumour cells. These latter are very uniformly dispersed and the majority are small. Morbid mononuclear and Hodgkin giant cells are identified in the mesh but most of the cells are slightly immature lymphocytes and their precursors. Eosinophils are present but not numerous. Plasma cells are noted but neutrophils are absent. Unequivocal mitotic figures are not numerous, average 2 per H.P. field; but densely staining nuclei are common. Necrosis is not present.

Diagnosis. Atypical Hodgkin's disease, very cellular, distinctly tumour-like.
This specimen is an example of reticulum cell sarcoma which shews mingled features of lymphosarcoma and true reticulum cell tumour. All vestige of normal structure in the affected node is lost. Where it is preserved the capsule shows marked fibrous thickening, but it is over-run and destroyed in most of the specimen. Fibrosis is present in the tumour but is relatively slight. Occasional perivascular lamellation and a little diffuse fibril formation are seen, several sharply demarcated areas of coralline deposit are noted. The predominant cell has the nucleus of a large immature lymphocyte which is rather pale staining. The cytoplasm is very indistinct. These cells are closely set in sheets but interspersed uniformly throughout them are, typical morbid mononuclear reticulum cells, Dorothy Reed mirror-image cells, and rare Hodgkin giant cells. Occasional cytoplasmic inclusions are visible in these cells possibly reflecting a macrophage activity. Eosinophilia is absent. Mitoses are numerous but necrosis is not present.

Diagnosis. Reticulum cell sarcoma, intermediate type.

This is a singularly fine example of early typical Hodgkin's disease. The new tissue appears in the form of indistinct nodules 0.5 – 2 mm. diameter which occupy about two thirds of the section. The remaining tissue is normal reactive lymphoid tissue, it is situated between the new nodules and at the sub-capsular periphery. The exodic growth phenomenon is well shewn and it is apparent that where the surviving lymphoid tissue is being crushed, the sinus tissue promptly disappears. In the field of the lesion fibrosis of all varieties is detected, in particular crescentic swathes formed by compaction of fibrils. The reticulum cell changes are typical in every way and Hodgkin giant cells are already numerous. Eosinophilia is pronounced. The mitoses vary locally but between one and two are observed in most high power fields.

Diagnosis. Hodgkin's disease, Typical. Early, still focal in the lymph node.
This node which is incomplete is entirely replaced by tumour. Low power examination conveys an impression of almost complete homogeneity but this is dispelled by high power study. The growth is a reticulum cell sarcoma which is shewing little tendency to sclerose. The cells are free spheroidal morbid reticulum cells in the looser parts of the tumour but their cytoplasmic outlines are not discernible in other areas. Cells corresponding to the large morbid mononuclear reticulum cell of Hodgkin's disease are very common and several almost typical Hodgkin giant cells are identified. The only detracting feature of the last named is their small size. Eosinophils are absent though plasma cells are present. Mitotic figures are very numerous, often over 20 being visible in a high power field. Superficially the lesion approximates closely to cellular Hodgkin's disease but the very inconspicuous fibrosis and lack of eosinophils removes it from this category.

Diagnosis. Reticulum cell sarcoma. Affinities to cellular Hodgkin's disease.
This specimen is remarkable. All the criteria for the diagnosis of Hodgkin's disease can be met, yet the picture is a quite astonishing parody of the typical disease.

The lymph node shows a fairly uniform pronounced fibrosis of the capsule and trabeculae. Diffuse fibril and fibrillary deposits, although tenuous, are present. Coralline deposits are absent, but perivascular lamellation is striking in some cases. The reticulum cell increase is almost entirely given over to the formation of large morbid mononuclear and Hodgkin giant cells. The former nearly all exhibit the "owl-eye" type of nucleus due to their pronounced nucleoli. Eosinophilia is emphatic and tends to be localised in the chinks between the large cell foci and the cells are set in among small lymphocytes. Plasma cells and neutrophils are scanty but recoverable. Mitotic figures are present in about average numbers; necrosis is limited to individual cells.

Diagnosis. Atypical Hodgkin's disease. Unusual type, overgrowth of morbid reticulum cells but with retention of typical complex.

This specimen is a lymphoid tissue tumour which does not conform to any orthodox category. It is nearest to a follicular lymphoma and has possibly evolved from this growth.

The capsule is thickened appreciably but is infiltrated and over-run in many places. Concentric re-formation by crescentic swathes of maturing fibrous tissue is notable. Trabeculae are represented, they shew irregular thickening and fuse with perifollicular fibrous tissue deep in the node. Fibrosis is expressed mainly in this fashion, but some perivascular lamellation and occasional coralline deposits are visible. The false follicles exhibit striking disparity in size and demarcation, many large ones appear compound due to subdivision, the nodules being delineated by surviving small lymphocytes. In places the pulp shews little follicular organisation and here morbid mononuclear reticulum cells are common. The pleiomorphism is not very pronounced and Hodgkin giant cells are rare. Eosinophilia is absent.

This is a follicular lymphoma with distinct affinity to lymphosarcoma. The capsule of the node is considerably attenuated and is over-run in the greater part of its extent. The tumour is composed of very numerous small solid type follicles which are chiefly composed of young lymphocytes, reticulum cells being unusually scanty. Fibrosis is very little in evidence, small fibrillary deposits are discernible in the remains of the pulp between contiguous follicles. Eosinophilia is absent. Reticulin impregnation confirms the follicular disposal of the tissue and so far as can be gauged there is little if any increase.

Diagnosis. Brill's disease, malignant. Affinity to small-cell lymphosarcoma.
This is Hodgkin's disease of an unusual type. The lesion is less pleomorphic than ordinarily and the reticulum cell proliferation is productive of very numerous macrophage type cells.

The specimen is a lymph node which is incompletely involved in the disease process. One pole shows well preserved lymphoid tissue in which the sinuses are packed with large macrophages. The node capsule is indifferently preserved but markedly thickened where present. The exocrine growth phenomenon is well shown. Fibrosis is pronounced. Perivascular and coralline deposits are present but not marked; the chief expression is symplasmic transformation, with the production of a uniform fine filigree. The reticulum cell proliferation already noted above does include infrequent morbid mononuclear cell types and very rare Hodgkin giant cells. Eosinophils are absent. The growth rate is apparently very slow.


This specimen is small, in fact fragmentary. It is described as tumour from the iliac fossa. It is apparently part of a lymph node which had previously been atrophic with fat replacement.

The lesion is probably a sclerosing reticulum cell sarcoma with morphological affinities to Hodgkin's disease but the specimen is inadequate for proper diagnosis. The reticulum cells are pleiomorphic and include many Hodgkin forms, eosinophils are present but scanty.

Diagnosis. Specimen inadequate. (Probably reticulum cell sarcoma, some affinities to Hodgkin's disease).

The specimen consists of two small abdominal lymph nodes replaced by reticulum cell sarcoma of intermediate (mixed) type with affinities to Hodgkin's disease.

Under very low power a distinct follicular arrangement is discernible. The tumour nodules are imperfectly outlined by swathes of fibrous tissue. The capsule is becoming thickened and is over-run at one point with new growth in the perinodal fat. Fibrosis is chiefly expressed as above, the centres of the nodules are almost purely nuclear. The type-cell has a large plump fairly pachychromatic nucleus with chromatin granules, but many typical morbid mononuclear reticulum cells are visible on close examination. Rare Hodgkin giant cells without typical cytoplasm are present. Eosinophils are present in sufficient numbers to be found easily. Mitoses are numerous up to 6 or 10 being countable in high power fields. Necrosis is not seen.

History. This patient had two previous biopsies on account of swollen lymph nodes in the axilla. The first is untraced, the second two months previously established a diagnosis of reticulo-sarcoma, the term being a generic usage. The third specimen M.R. 540:41 consists of four small axillary nodes and the scar of the previous biopsy. The fourth biopsy is a nodule removed from the abdominal wall several months later.

One of the axillary nodes in hyperplastic and seen alone could not be regarded as neoplastic, incipient abnormality is restricted to loss of sinus tissue and slight immaturity of the lymphocytes. The remainder are replaced by active lymphoid tissue tumour which is characteristically homogeneous. The capsules are of normal thickness, hyaline and at several places are over-run by tumour cells. Invasion of adjacent fat is conspicuous. The type-cell stains almost as deeply basophil as an immature lymphocyte but is larger and has often the misshapen yet distinct nuclear membrane of a morbid reticulum cell. Cytoplasm is impossible to define in most of these cells. Typical morbid mononuclear reticulum cells are present, but Hodgkin giant cells are not. (Foreign body giant cells with inclusion are numerous in the scar). Occasional eosinophils are seen but not plasma cells. Mitoses are numerous and necrosis, especially pyknosis, is pronounced. There is a conspicuous lack of fibrosis, a little fibril formation is seen but that is all. In the abdominal deposit isolated fragments of skeletal muscle simulate coralline fibrosis.

Diagnosis. Lymphoid tissue sarcoma. Intermediate type (Lympho Reticulo-sarcoma), Lymphoid tendency preponderating.
History. Unknown beyond the fact that the specimen is a cervical lymph node which had been enlarged (solitary) for about a year.

Superficially the appearances are typical follicular lymphoma. Scrutiny reveals that some of the peripheral follicles have normal cells in their pale centres and are surrounded by thin imperfect zones of normal small lymphocytes. The false follicles display the usual low power appearance of homogeneity.

The capsule is greatly thickened, being over 1 mm. deep in places, it is densely hyaline but tumour cells are insinuated into chinks in it. The exodic growth phenomenon and concentric re-formation of capsule are evident. True trabeculae are scanty but new swathes of connective tissue are conspicuous between the follicles. The fibrosis is unusually copious in this case; perivascular lamellation, fibrils, and even coralline deposits are seen. The tumour cell type tends to have a reticulum cell type of nucleus but it is small. A very few small atypical Hodgkin giant cells are present. Eosinophils, plasma cells, and neutrophils are inconspicuous. Growth rate is slow and necrosis is not a feature.

(i) Tonsil. (ii) Recurrence in Uvula two years later (reference relates to second specimen.

The specimen of palatine tonsil shows early neoplastic change in the pulp lymphoid tissue. The growth is a uniform cellular tumour of low mitotic activity. The type cell has a reticulum cell nucleus but the cytoplasm stains similarly to plasma cell cytoplasm. Fibre formation is little in evidence. Numerous apparently normal follicles of normal-sized small lymphocytes are scattered through the growth. In some of these follicles small pale centres are visible. Polymorphs are present in the crypts but in much the same quantity as in normal tonsil.

In the recurrence specimen the malignancy of the tumour is enhanced, and fibrosis is more in evidence. Mitoses are numerous and the previous uniformity of cell type is being lost. Occasional multinucleate giant cells are identified which are smaller than typical Hodgkin giant cells but otherwise closely similar.

Diagnosis. Reticulum cell sarcoma, sclerosing slightly. Becoming pleiomorphic.

The specimens are unfortunately not good, fixation and cutting are poor. The lesion present is atypical Hodgkin's disease which has a tumour-like homogeneity and peculiar giant cells.

Traces of surviving normal lymphoid tissue can be identified; they include fragments of marginal sinus tissue and peripheral follicles of solid type. The capsule shows only doubtful thickening. Trabeculae are not represented. Fibrosis is curiously patchy; perivascular lamellation and capillary hyalinisation are notable. Diffuse fibril, fibrillary, and coralline deposits are also present. The reticulum cell proliferation has not the usual capricious heterogeneity and the commonest cell-type present is lymphoblastic in type. Morbid mononuclear reticulum cells are present, and typical Hodgkin giant cells can be seen. Small multinucleate giant cells which are nearer to the foreign body variety than to Langhan's type are conspicuous, in some cases these are huddled in little clusters round coralline collagen deposits. Eosinophilia is present but is of a low order. Other cells are scanty. Mitoses are identified with difficulty in the preparations.

Case of Cleminson. Ref. 213:38. Cervical lymph node. History unknown beyond the fact that this patient was recorded to be alive and well three years later. Provisionally labelled by J.S.D. as "? reticulosis, with uniform small cells, fibrosis".

The specimen is an enlarged incomplete lymph node. Where preserved, the capsule is uniformly hyalinised and just appreciably thickened. Several crescentic bands of hyaline connective tissue are visible which segment the node imperfectly into nodules 2-3 mm. diameter. Low power the lymphoid tissue appears replaced by a uniform small cell growth in which a few, small, widely dispersed germinal centres survive. These have tiny irregular pale centres, 7-10 cells across, in which the component cells have pleiomorphic nuclei but few or no mitotic figures. The peripheral lymphocytes are typical small lymphocytes. In comparison the type-cell nucleus of the diffuse growth is larger and leptochromatic - a less mature lymphocyte. In addition to these cells slightly enlarged reticulum cell nuclei are apparent but typical Hodgkin giant cells are not present in any of the material available. A few eosinophils are seen near the fibrous septa, other cells are conspicuously absent. The growth rate is low in spite of high cellularity, there is seldom more than one mitotic figure in high power fields. (Many blood capillary vessels with much swollen endothelial cell nuclei present a similarity to giant cells but their true nature is clear on scrutiny).

Diagnosis. Benign Hodgkin's disease.
Mr. D. Presented initially with lymph node enlargement in the neck stated to have been present for ten years. Biopsy was performed 21st May 1936 ref. P.S. 87:36. A further microscopical examination ref. P.S. 5:37 was made 7th January 1937 after irradiation. The patient was reported alive and well in December 1938.

The first specimen is a lymph node in which the capsule is slightly thickened and densely hyaline except at a few sites where its composite lamellae are split by infiltrating lymphocytes. This invasion is possibly artefact and is distinctly limited. Survival of normal tissue is minimal and takes a curious form; there is a thin but definite peri-thelial cuff of normal small lymphocytes round most of the small blood vessels deep in the node. These serve to emphasize an otherwise very indistinct false follicle arrangement of the tumour tissue. The tissue is composed of mutually identical cells which correspond to rather young lymphocytes, even high power the purity of the cell type is remarkable. Fibrosis is present; for the greater part it consists of scar-like deposits of the coralline variety which are very similar to those observed in Hodgkin's disease. A small contribution of fibrils and perivascular lamellae make up the remainder. A very few morbid mononuclear cells are present but there are not any giant cells. Eosinophils, plasma cells and neutrophils are exceedingly rare. Mitoses are scanty and necrosis is not seen. The second specimen shews what is virtually an infarcted node, there is a large central area of coagulative necrosis. Adjacent to this is a young connective tissue mantle with bizarre reticulum cell or fibroblast nuclei and moderate eosinophilia. Curiously, the arterioles do
not shew any endarteritis.

The specimen represents part of a large fused lymphoid tissue mass. Except at one edge where fragments of the original capsule persist all vestige of normal structure has gone. The capsule is grossly thickened and hyaline, and it is continuous with a very thick fibrous mesh in which are inserted numerous small nodules of typical Hodgkin tissue 1-4 mm. diameter. Most of the fibrosis takes the form of mature swathes of compacted fibres, either fibrillary or coralline deposit is evident. The islets of cellular tissue are composed of young lymphocytes, morbid reticulum cells including giant cells, eosinophils, and fibroblasts. Transitions between reticulum cells and fibroblasts are common. The eosinophilia is unusually copious considering this is a late stage. Necrosis with many pyknotic polymorphs is noted in many foci. Plasma cells are rather remarkably confined to the least mature (innermost zone) connective tissue fringing the follicles. In a slide stained specifically for eosinophils there is very little evidence of transformation of plasma cells to granular series. Mitoses average 1 per high power field. The vascularity of the lesion is low; vessels of arteriole size are rare.

Diagnosis. Hodgkin's disease, Typical, late, shewing very striking follicular pattern.

History. A painless swelling appeared in the left side of the neck about five months ago. An initial clinical diagnosis of tuberculous lymphadenitis was abandoned since the tumour increased steadily in size. The patient has no other complaints.

This is a cervical lymph node shewing an unusual form of atypical Hodgkin's disease. The reticulum cell proliferation tends to be focal and eosinophils are few. A small adjacent node included in the specimen is still physiological. There is some persistence of normal lymphoid tissue in the larger (morbid) node. This appears as small crushed crescent shaped deposits being driven against the capsule. The exodic growth phenomenon is well illustrated. The habit of growth is peculiar, all elements of the Hodgkin's disease complex are discernible but the reticulum cell proliferation is focal, large masses of morbid mononuclear and Hodgkin giant cells forming irregular islets and strands. The capsule is thickened and hyaline, and concentric re-formation is visible. It is not infiltrated.

Fibrosis is seen in all forms except coralline deposits. The reticulum cell proliferation appears excessive due to the fact that the cells are not uniformly disseminated. In places they are symplasmic. Eosinophils are present but in small numbers. Plasma cells and neutrophils are present but meagre. Mitoses average 1 per H.P. field. Necrosis is not a feature.

Diagnosis. Atypical Hodgkin's disease. Eosinophils few. Reticulum cell proliferation is tending to be focal.

Note. This patient died in 1951. Full necropsy was carried out see special P.M. series, E.35:51.
Both specimens shew the same features. The lesion does not conform to any of the commoner variants of lymphoid tissue tumour but is nearest to follicular lymphoma.

Fibrosis is a well developed change. The capsules are considerably thickened and the trabeculae are similarly affected. In addition perivascular lamellation is notable in the trabecular vessels. Fibrils and fibrillae are present in the pulp in relation to macrophage foci and near the edges of follicles. Follicles of small size and irregular shape are present throughout the nodes, they are mainly of the solid variety and the centres are composed of active reticulum cells differentiating to lymphoid type, mutual compression of follicles is seen in some areas. The pulp contains many aberrant cells of morbid mononuclear reticulum cell-type. Rare Hodgkin giant cells are identified. Eosinophilia is absent.

In both nodes large focal areas of necrosis are present and some disintegrated polymorphonuclear leucocytes are visible at their periphery. In some areas aggregates of macrophages are seen, many of these cells are degenerate but between the surviving ones a fine collagen mesh is discernible.

Diagnosis. Lymphoid tissue tumour, indeterminate, nearest to Brill's disease. Focal necrosis, fibrosis and abnormal proliferation of cells in the node pulp.

This specimen consists of skin and subcutaneous tissue.  In the later coalescent deposits of Hodgkin's tissue are present together with physiological granulation tissue and purulent abscess. The Hodgkin's tissue contains numerous evenly dispersed morbid mononuclear reticulum cells and a small number of typical Hodgkin giant cells. Lymphocytes and eosinophils are copious, many of the latter are mononuclear and of lymphocytoid type. The granulation tissue contains fairly numerous iron pigment-laden histiocytes. These cells are conspicuously different from any in the tumour proper, the nuclei are typically small and eccentrically placed. Fibrosis is difficult to assess due to pre-existing collagen in the site, but a fine fibrillar mesh is discernible in the tumour and is continuously related to morbid reticulum cell processes which taper off into fibrillae. The morbid process is readily recognisable as Hodgkin's disease despite its superficial uniformity. The abscesses noted shew central necrosis and abundant polymorphs, capillaries are little in evidence. Localisation of the lesion is sharp.

Unknown. Identity ref. 113242.

The specimen consists of four lymph nodes, two moderately enlarged, the others of normal size. All shew uniform but slight thickening of the capsules, in one case two adnate nodes shew mutual destruction of their capsules at the region of contact. In most areas there is a very thin rind of compressed but otherwise apparently normal lymphoid tissue including rare fragments of sinus tissue. This exodic growth manifestation is disappearing in places. Trabeculae are few, those present shew incipient accretion of collagen fibres along their course but these are not yet compacted. Fibrosis is less than usual in typical Hodgkin's disease; perivascular lamellation, diffuse fibril deposition, and symplasmic transformation are the main expressions of it. Coralline deposits are not seen. The reticulum cell increase is well above the usual and appears much at the expense of lymphocytes. Lymphocytoid differentiation is common, but typical morbid mononuclear type reticulum cells are more numerous. Hodgkin giant cells are present but tend to be small with few nuclei. Eosinophils are numerous, unexpectedly so for this cellular type of lesion. Plasma cells and neutrophils are also commoner than usual. Mitoses are up to 4 per H.P. field. Necrosis, except for occasional individual cells, is not a feature.

Diagnosis. Hodgkin's disease, tumour-like type.
The specimen is a lymph node entirely replaced by reticulum cell sarcoma. There is not any vestige of normal structure except for portions of the original capsule. This tunic is thickened and hyaline where it persists and concentric re-formation is noted. Elsewhere it is over-run and destroyed by the tumour cells which infiltrate out into the adjacent areolar tissue. True trabeculae are not represented and the new collagen of the tumour is still diffuse. Perivascular lamellation is not a feature, the fibrosis is chiefly expressed by diffuse fibre and fibril formation which imparts an indistinct alveolar pattern to the tumour. Coralline deposits are common and occasionally appear to be condensations of the ground fibres. The type cell of the growth is a reticulum cell with a rather small nucleus, superficially it resembles a lymphocyte series cell but high power scrutiny negatives this. Nucleoli are common and typical morbid mononuclear reticulum cells are identified readily. Hodgkin giant cells are present but forms with more than three nuclei are rare. Eosinophilia is absent and plasma cells and neutrophils are inconspicuous. The growth rate is high judged by Karyokinetic figures which average about 4 per high power field.

Diagnosis. Reticulum cell sarcoma, sclerosing. Fields of Hodgkin's disease are present (less topical eosinophilia).
62.

Ref. R.M. Cervical lymph nodes.

This is atypical Hodgkin's disease which is very tumour-like in appearance and is beginning to conform to Warthin's Hodgkin's sarcoma, type II (Reticulo-endothelioblastoma). Normal lymphoid tissue does not persist except for small islets at the edge of the nodes. The capsule is markedly thickened and extremely hyaline. There is much fibrosis in the node. Most of it assumes the form of irregular crescentic swathes and small irregular stellate scars; under low power an interlacing of the fibres, reminiscent of fibro-sarcoma, is appreciable. Perivascular deposit is common too, but coralline formations are scanty. The reticulum cell increase is unusually florid and both morbid mononuclear and Hodgkin giant cells are particularly numerous. Eosinophilia is present but rather below average intensity. Mitotic activity is very difficult to assess, there are so many aberrant hyperchromic nuclei present; unequivocal examples of Karyokinesis are rare. Several small necrotic foci are present.


This is an example of soft cellular rather tumour-like Hodgkin's disease. Polymorphonuclear leucocytes are profusely scattered through the new tissue. Small poorly demarcated islets of lymphoid tissue and fragments of marginal sinus survive at some points of the periphery. The capsule is thickened but is still cellular, trabeculae are not represented. Fibrosis is slight, all varieties are encountered but most is fibrillary. It forms a fine filigree and is a good example of symplasmic transformation. Almost all the fibres stain red with van Gieson's stain. Reticulum cell proliferation is pronounced and lymphocytes are very much repressed. Most of these reticulum cells are small and uniform though typical morbid mononuclear and Hodgkin giant cells are present. Eosinophilia is marked despite the large number of neutrophils present. Small necrotic patches are frequent. Under very low power the exodic growth phenomenon is appreciable at some places.

This is typical Hodgkin's disease, the lesion is fully established; the new connective tissue renders the cellular part of the node folliculoid. There is not any persistence of normal lymphoid tissue. The capsule is grossly thickened and hyaline with numerous trabeculae running from it to cut the node into a mosaic. In addition to this coarse fibrosis, perivascular deposits and fibril formation in the cellular islets are common. Coralline formation apparently the result of compaction are also found. Reticulum cell proliferation is pronounced in many of the islets with copious production of morbid mononuclear and Hodgkin giant cells. The latter are frequently necrotic and they are small. Eosinophils are numerous, often focal, and mononuclear forms are common. Plasma cells and neutrophils are present in about the usual concentration. Mitotic figures are difficult to identify with certainty but are probably few. Several minute fibrinoid necrotic foci are present.

Diagnosis. Hodgkin's disease, Typical.
Ref. G.R.I. 1168:43. A large single lymph node 4 cm. diameter.

This is classical Hodgkin's disease at a moderately early stage. The entire node is morbid, the capsule is fairly evenly thickened being approximately 1 mm. wide but trabeculae are hardly represented. Incipient concentric re-formation of the capsule is indicated but the true capsule is devoid of any infiltration. Fibrosis of all varieties is encountered; perivascular lamellation is present throughout and swathes of new fibrous tissue are being compacted into bands which are forming the new tissue into nodules. Fine fibrillar filigrees are formed in the more cellular areas and some coralline fibrosis is observed. The reticulum cell proliferation is general and besides the differentiation (or reversion) to morbid mononuclear and Hodgkin giant cells there is easily traced transformation to fibroblast and epithelioid type cells. A few unusual giant cells are noted, their nuclei shew a horse-shoe arrangement but are otherwise unlike Langhan's cells. Eosinophilia is marked, plasma cells and neutrophils are present in small numbers. Mitoses average 1 per H.P. field; necrosis is limited to individual cells.

Diagnosis. Hodgkin's disease, Typical. (Moderately early).
This is a good example of typical Hodgkin's disease, the eosinophilia is unusual in its intensity.

Several survival follicles with pale centres and survival islets are present, they are situated relatively deep in the node. They appear remarkably incongruous being surrounded by typical Hodgkin's tissue. The capsule is grossly thickened and hyalinised. New connective tissue trabeculae of very irregular size and course are seen. All forms of fibrosis are present in quantity, the change is coarse. The reticulum cell proliferation presents in the usual degree; fibroblastic transformation is well shewn besides the characteristic morbid mononuclear and Hodgkin giant cells. The latter are typical but necrosis is common in them. The eosinophilia is a most impressive element in this case. Some high power fields seem almost totally composed of these cells. Plasma cells and neutrophils are present but in the usual quantity. Mitoses number 1-2 per H.P. field. Focal necrosis is common, the demarcation of these areas is curiously sharp.

Diagnosis. Hodgkin's disease, Typical. (Eosinophilia very pronounced).

This is atypical Hodgkin's disease of tumour-like type. Eosinophils are almost absent and the growth is relatively homogeneous with a preponderance of small cells. The capsule is grossly thickened and islets of the new tissue are set in it, this phenomenon appears the result of concentric re-formation of the capsule with engulfment of the cellular parts, rather than transgression of the capsule. Fibrosis of other types is much less in evidence. The reticulum cell proliferation is productive of young lymphocyte type cells, the typical mononuclear and Hodgkin giant cells being overshadowed. Eosinophils are very rare, but plasma cells and neutrophils are above average in number. Mitoses are common, up to 4 per H.P. field. Necrosis is confined to individual pathological cells.

While areas of typical Hodgkin's disease are easily found the greater part of the tissue is not orthodox.


This is an example of atypical Hodgkin's disease of an unusual and somewhat treacherous character. The true nature of the change is obscured by the development of epithelioid cells which occur in irregular patches. High power scrutiny dispels all doubt of the true pathology.

The exodic growth phenomenon is present but is indistinct. The peripheral rind of tissue is hardly normal but contains many more small lymphocytes than the new tissue. The capsule shews local thickening and concentric re-formation is discernible. Fibrosis of all varieties is identified but it is small in amount and difficult to find without special staining. The reticulum cell proliferation is expressed in the usual way with perfectly typical morbid mononuclear and Hodgkin giant cells but epithelioid cells are much more conspicuous. They are disposed in irregular patches without any tendencies like 'endothelial tuberculosis'. Eosinophilia is moderate, plasma cells and neutrophils are very scanty. Mitoses are numerous for Hodgkin's disease, 2-3 being visible in certain H.P. fields; two minute areas of necrosis are seen.


This is early Hodgkin's disease. Most of the capsule is lost by artefact and it is not possible to gauge whether thickening is present. Fibrosis is already fairly pronounced all forms being encountered but coralline deposit is very scanty. The reticulum cell hyperplasia is only moderate. Most of the cells produced appear to be immature lymphocytes but the pathognomonic ones are also present. Eosinophilia is confirmed; the staining leaves much to be desired but their number is probably high. Plasma cells are scanty but neutrophils are scattered diffusely through the tissue. Mitoses are numerous, 4-5 per H.P. field. Necrosis is limited to individual cells.

(Very little material is available from this case, the spare blocks having been used for class purposes).

Diagnosis. Hodgkin's disease, Typical, Early.
This is a good example of early, established Hodgkin's disease which is not involving the whole node and demonstrates the exodic growth phenomenon. The new tissue occupies the central part of the node and is growing just like a metastatic cancer nodule and driving a rind of reactive lymphoid tissue up against the capsule. The capsule is not thickened but the marginal sinus is obliterated, at the margin of the tumour tissue there is a fibrous layer forming, an example of early concentric re-formation of the capsule. Fibrosis of all varieties is already evident. The reticulum cell increase is of the usual order and productive of typical morbid mononuclear and Hodgkin giant cells. Eosinophils are numerous. Plasma cells and neutrophils are little in evidence. Mitoses average about 1-2 per H.P. field. Necrosis is not seen.


Swelling in the left side of the neck of several months duration.

This is early Hodgkin's disease. There is remarkable survival of physiological lymphoid tissue. Follicles, both solid and with pale centres, are present. The latter are restricted to the periphery of the node. Fragments of sinus tissue are also discernible. The pale centre reticulum cells do not appear to contribute to the new tissue which in this case is convincingly restricted to the pulp (functional medulla of Robb-Smith).

The capsule shews irregular thickening, most of it being seen at the base of small wedge-like trabeculae. The exodic growth phenomenon is well demonstrated and concentric re-formation of the capsule is striking. An inner false capsule round the outer margin of the new tissue is distinctly visible. Fibrosis is well established, all forms are identified but the coralline variety is minimal. Reticulum cell proliferation is conspicuous and morbid mononuclear types are very numerous. Typical Hodgkin giant cells are common. Several Langhan's multinucleate giant cells are present and appear directly related to foci of necrosis. Eosinophils are very numerous, several large foci of several hundred are seen. Mitoses number approximately 2 per H.P. field. Necrotic foci are common, they are chiefly at the expanding edge of the lesion. (See also Serial 76).


Specimen consists of four large smooth nodes, the largest 4 cm. diameter.

This is early Hodgkin's disease. The coarsely follicular organisation of the new tissue is definite and the exodic growth phenomenon together with concentric re-formation of the capsule is well shewn. The tumour is atypical, small cells being predominant and eosinophilia slight.

There is patchy survival of lymphoid tissue and sinus tissue at the periphery of some of the nodes. The capsule is distinctly thickened, this is uniform but only moderate in degree. Fibrosis of all varieties is encountered but is slight overall. The reticulum cell proliferation is less prominent than usual due to the high number of small and slightly immature lymphocytes. Morbid mononuclear and Hodgkin giant cells are identified but they are scanty. Eosinophils require a patient search, they are present but distinctly rare. Mitoses are few, less than one can be found consistently in high power fields. Necrosis is absent.

Diagnosis. Atypical Hodgkin's disease. Small cells predominate, no eosinophilia, fibrosis slight. Some affinities to small cell lymphosarcoma.

Note: Patient died 2nd December 1950, P.M. 332:50. G.R.I. q.v.
Thomas Finnigan.  Aet 74 years.  G.R.I. 2387:47.  Enlarged axillary lymph nodes for doubtful period, probably months.  Also has tumour over frontal bone and in chest wall.  Specimen of axillary node sent.  See also serial 135.

This is a poorly differentiated reticulum cell sarcoma which exhibits some affinity to Hodgkin's disease.  The entire node is neoplastic, the capsule being extensively destroyed and the tumour invading the surrounding fat.  The growth assumes the general form of alveolar sarcoma, groups of cells occupying the spaces of a plexiform fibrous tissue mesh.  The cell type is spheroidal with indistinct cytoplasm of pale amphophilic staining reaction and morbid reticulum cell nuclei of various shapes and sizes.  The cells are discrete in some areas and fused to form plasmodial masses in others.  Typical Hodgkin giant cells are present.  Eosinophilia is absent and plasma cells are not apparent.  Polymorphs are notable round small foci of necrosis which are seen.  Perivascular fibrosis is seen round the arterioles present in the tumour.  The activity of the growth is high, 4-5 mitotic figures can be found in most high power fields.


This is an example of moderately early Hodgkin's disease in which the origin of some of the integral fibrosis is well seen.

The node capsule is of normal thickness, it is intact and concentric re-formation is not evident. Several ill defined pre-existing lymphoid follicles are discernible. Fibrosis is slight but well beyond the amount seen in normal nodes. Perivascular lamellation and fine fibrillary deposits are the chief expression. The latter is traceable to the cytoplasm of reticulum cells which have assumed an epithelioid appearance. While the nuclei of these cells is close to those of proliferating fibroblasts the cytoplasm is rarely spindle-shaped. These fibrifying reticulum cells are the commonest of this scion but morbid mononuclear and Hodgkin giant cells are plenteous. A few of the latter shew horse-shoe arrangement of their nuclei. Eosinophilia is conspicuous but the other cells, plasma cells and neutrophils are scanty. Mitoses average 1 per H.P. field, necrosis is absent.

Diagnosis. Hodgkin's disease, Typical, early, established. Diffuse involvement of almost whole node structure.
Isobel Topping. Aet 36 years. Under Dr. David Smith. Ref. G.R.I. 110:49. Complaint of malaise and swelling of the right side of the neck for five weeks. Right tonsil enlarged. Firm discrete glands in the right posterior triangle of the neck. The peripheral blood is normal. (This patient later developed Hodgkin's disease in the breast, see Serial 81. (Ref. G.R.I. 2583:50) Known to be dead, 1952, necropsy not performed.

This is early Hodgkin's disease. The entire node appears affected but small lymphocytes are still copious, these particular cells are just appreciably less mature than normal ones. The capsule already shews hyaline thickening and fibrosis of all types is present. Perivascula r deposit is the chief expression of the change. The reticulum cells shew a very uniform increase, many morbid mononuclear forms are present but in general they are small. Typical large ones and Hodgkin giant cells are however present. Eosinophilia is less conspicuous than usual and tends to be focal. Plasma cells and neutrophils are scanty. Necrosis is limited to occasional giant cells. (Note, the peculiar veins of Ehrich are packed with normal small lymphocytes).

Diagnosis. Hodgkin's disease, Typical, early, established.

The specimen is just recognisable as a lymphoid structure. The capsule is greatly thickened by lamellae of collagen but nuclei of fibroblast and fibrocytes can still be made out. At one point a portion of the marginal sinus persists. Areas of the characteristic new tissue are recognisable but certain changes are noteworthy. Plasma cells, frequently binucleate, are common, and the fibrosing process is florid. Eosinophils are absent but occasional polymorphs are seen. Numerous pyknotic nuclei are scattered widely through the tissue, these appear to be chiefly lymphocytic.

Enhancement of fibrosis, loss of eosinophilia and abundance of plasma cells appear to be the chief features.

Diagnosis. Hodgkin's disease, post-irradiation.
Complaint of sudden severe swelling of all limbs which persisted for four weeks and then subsided. Enlarged cervical nodes for past week. Exceedingly ill; the peripheral blood is normal. The biopsies are of cervical and epitrochlear nodes and were taken at an interval of six weeks. The patient died six weeks after the second operation, necropsy was not obtained.

The microscopical appearances are the same in both specimens apart from slight increase in the fibrous tissue of the capsule in the second node. This is cellular rapidly growing Hodgkin's disease which clinically merits the term acute Hodgkin's disease. Persistence of normal structure is seen, in the first node several lymphoid follicles persist and in the second much of the marginal sinus remains but the lumen contains abnormal cells.

Capsular thickening is seen only in the second specimen. This barrier is not invaded but small islets of the new tissue, with many plasma cells, are enclosed in the substance. Fibrosis is relatively slight; perivascular lamellation is certainly present in the few vessels of size present, but diffuse fibril and fibrillary deposits are the chief form. Very early coralline deposit is shown in process of forming and the node of origin is beautifully disclosed. A focus of morbid reticulum cells which have assumed an epithelioid morphology are beginning to fibrose, the process being a direct transformation of their cytoplasm into collagen, this commences as a condensation of their peripheral cytoplasm into
into fibrils which by mutual accretion form the familiar tortuous cylinder. The reticulum cell proliferation is pronounced and many typical morbid mononuclear forms are present. Hodgkin giant cells are present, they tend to be rather small. A few giant-cells shew similarity to Langhan's type but their nuclei are not characteristic. Eosinophils though present are scanty, similarly the plasma cells and neutrophils are few. Mitoses average 4-5 per H.P. field.

Mrs. Caroline Cameron. Aet 42 years. Ref. 2444:49. Patient was admitted for investigation of suspected intestinal obstruction. Laparotomy was carried out 26th October 1949 and an inoperable retroperitoneal tumour was found. This specimen consists of a portion of this growth. The patient died three days later. An incomplete necropsy was performed but, there is not any record available. (From an outside hospital, without a morbid anatomist).

The specimen shews a limiting cellular membrane at one edge which is serosa, beneath this a zone of fatty areolar tissue overlies an highly cellular tumour which is widely necrotic.

The growth is morphologically a pleomorphic reticulum cell sarcoma in which morbid mononuclear type and Hodgkin giant cells are present. Eosinophils are very numerous; plasma cells and neutrophils are copious near the necrotic foci. Mitotic activity is considerable. Fibrosis and new reticulin formation are confirmed by special staining.

The special interest in this case lies in the remarkable similarity of certain areas of the tumour to Hodgkin's disease. Morphologically it is a link between the two, its lymphoid origin is probable as in some sections fragments of lymph node are visible. (The necropsy sections shew deposits of the same tumour in the bowel).

Diagnosis. Reticulum cell sarcoma, pleiomorphic, Eosinophilia present.
British Soldier. Aet 23 years. Ref. L.N. 20:50. The specimen is a cervical lymph node received from Capt. Cowan, R.A.M.C., Cowglen Military Hospital. Complaint of swelling in the neck, left side, of uncertain duration. The general health of the patient is excellent.

The capsule shews irregular thickening in places. Where it is still normal there is surviving sinus tissue and a thin rind of subjacent lymphoid tissue. These normal tissues are hyperplastic but still in physiological degree. The exodic growth phenomenon of the tumour is particularly well shewn, and capsular re-formation is seen, sometimes outflanking survival islets of non-neoplastic lymphoid tissue. The pre-existing trabeculae are thickened and new swathes of connective tissue are forming. Fibrosis is absolutely typical; perivascular lamellation, fibrils, fibrillae, and coralline deposits are all seen. The reticulum cells are conspicuously increased and the majority are pleiomorphic morbid mononuclear types. Hodgkin giant cells are prominent. Topical eosinophilia is marked; plasma cells and neutrophils are also seen. On an average two mitoses are seen in high power fields. Minute foci of necrosis are notable. (This is amongst the most classical examples of the disease at an established yet early stage that I have seen).

Diagnosis. Hodgkin's disease, Typical, early, established.

History. Patient has noticed glandular swellings in his right elbow region and the right axilla for three years. Anaemia is present. (The first biopsy was mis-diagnosed as reactive). At the second operation the axilla was cleared entirely, with apparent benefit. He reported back in July 1950, but did not return. He died 22nd December 1950, at home, necropsy was not obtained.

This is an example of an unusual variant of Hodgkin's disease. It corresponds most nearly to Ziegler's Type II, but does not conform exactly.

The capsule and trabeculae are thickened sufficiently to arrest attention. The lymphoid tissue of the node is rendered into small pseudo-follicles in most areas, by incipient perivascular fibrosis of the peculiar veins of Erich. The tissue thus arranged looks superficially normal but close scrutiny reveals quite numerous pathological reticulum cells and rare Hodgkin giant cells. The lymphocyte is still by far the commonest cell present but it is an immature type with a large nucleus which is relatively leptochromatic. Fibrosis of fibril and fibrillary type is discernible, especially in the second specimen, but it is not obtrusive. Eosinophils are absent from both. Plasma cells are present in larger numbers than is usual in the disease. Mitoses are very infrequent, many fields being devoid of them.


History. This patient came under the care of Mr. Stevenson in July 1950 on account of a localised swelling in the breast. This tumour was resected and sent for microscopical examination. At this time the previous history (See Serial 75) was unknown.

The specimen consisted of a firm fibro-fatty mass rather larger than a walnut.

Breast tissue of normal appearance is readily identified in parts of the specimen. The normal connective tissue of the gland, both the primitive gland field variety and the general stromal type present usual appearances. Several small and one large (1 cm. diameter) deposit of typical Hodgkin's disease are present; there is not any evidence that these have arisen in ectopic lymph nodes. The tumour shews well developed fibrosis which is coarse and hyaline, and differs sharply from the native formations. The cellular component of the lesion is quite characteristic and is some areas typical Hodgkin giant cells are very numerous. Eosinophilia is slight. A distinct irregular follicular growth is notable. In places surviving epithelial structures, acini and ducts, are over-run by the growth.


History. Patient complained of a swelling in the left inguinal region which had originated four years previously. This increased in size rapidly during the last two months. Aspiration was attempted in November 1949. The present specimen was resected 26th November 1950.

The specimen is presumed to be a part of an enlarged lymph node but vestiges of original structure are absent. This is an example of reticulum cell sarcoma corresponding exactly with the clasmatocytic stem cell lymphoma. The type cell is a morbid mononuclear reticulum cell; many shew acidophile nucleoli. In all areas the cells are mutually separated; several typical Hodgkin giant cells are present. Fibrosis is seen chiefly as perivascular lamellae and scar-like areas of compacted fibrils. In addition a fine filigree of stout reticulin fibres is demonstrable. Eosinophils are very few and a few polymorphonuclear leucocytes are seen near necrotic areas. The latter are massive infarcts. Mitoses are very numerous, about 20 per high power field.


Note. This patient died a few weeks later from pulmonary embolism, and necropsy (G.R.I. P.M.304:50) revealed restricted spread of the disease, in the abdomen and thorax. Deposits were confirmed in the liver and in abdominal nodes by microscopy.

History. This patient was unable to offer any account of his illness, his cerebration was very slow. The presenting symptoms were enlargement of the axillary lymph nodes and jaundice. The initial biopsy, (763), was from the left axilla, the second, (932), from the portal fissure.

This is an example of classical Hodgkin's disease which was originally missed. The error was made because the first sections shewed a lymph node in which fibrosis was the only morbid feature of note. Further sections revealed the true character of the lesion. This case emphasizes the paramount importance of appreciating that fibrosis of obscure cause in lymph nodes is a most sinister finding. In both specimens surviving lymphoid tissue, including follicles, pulp, and sinus tissue, persist. The exoctic growth phenomenon however is not displayed, the growth of this lesion is relatively indolent. The capsules shew irregular fibrous thickening and trabeculae are visible. Fibrosis of all varieties is encountered but by far the greatest portion is coralline. Morbid mononuclear cells, Dorothy Reed mirror-immage cells, and Hodgkin giant cells are all present. Eosinophils are present, but slightly below the usual number. Even in the cellular areas of the lesion small lymphocytes are conspicuously few.

Diagnosis. Hodgkin's disease, Typical, late.

History. Patient complained of a small lump present behind the left ear which was present two months. During the past week another swelling has started below her chin. Pediculosis capitis was present. After biopsy the wound healed rapidly and the patient was dismissed with instructions to return in three months. She has never returned to date (November 1954).

This is a slightly enlarged lymph node showing pronounced reactive change in which epithelioid cell change is seen throughout the pulp. In some areas the appearances very closely simulate Hodgkin's disease but the reacting cells are distinctly small and the general architecture of the node is retained. Fibrosis is present but is not of Hodgkin's disease conformation.

There is general retention of normal architecture, this is difficult to appreciate with the ordinary low power but is well seen with lower magnification (X 30). The germinal follicles are irregular in outline with occasional abstraction of fragments of the pale centre tissue. The pulp is mottled due to numerous small aggregates of epithelioid cells and rare individual large reticulum cells. The capsule is thickened and the fibres are hyaline; in most parts a fairly uniform infiltration of small lymphocytes and plasma cells is present. Fibrosis is seen in the trabecular vessels, and round other vessels in the node. Very fine fibrillae of collagen are present in the cytoplasm of many of the epithelioid cell collections. The reticulum cells encountered have been noted.
noted above, very few shew hyperchromatic or distorted nuclei. Rare mirror-image cells are seen but again they appear physiological. Hodgkin giant cells are not present. Eosinophils and polymorphs are present in small numbers. Mitoses are chiefly seen in the pale centres but several are also noted in the pulp.


History. Painless swelling in right side of neck of three months duration, latterly swelling has appeared in the right axilla. At operation two masses of cervical lymph nodes were removed one weighing 70 gm. and the other 15 gm.

This is a very interesting lymphoid tissue tumour with a peculiar morphology. Features of Hodgkin's disease and large cell lymphosarcoma are intermingled giving what may be described as "lymphoblastic Hodgkin's disease". There is very little remaining normal lymphadenoid structure, survival islets of small lymphocytes constitute the sole evidence. The capsule is irregularly thickened and hyaline, slight concentric reformation is evident and invasion is present. Trabeculae are few but partake of the capsular thickening. Fibrosis is present but slight. Patches of diffuse fibrils are seen together with very delicate sinuous fibrillae and some typical coralline deposits are seen. The majority of the cells are mutually similar. They are free cells with round nuclei, folds and wrinkles of the fairly dense nuclear membranes are rare. Chromatin content is variable but usually large, in the forms of indistinct blobs. Amphophilic nucleoli are very frequent. The cytoplasm is oxyphile, very finely granular, and concentric round the nuclei; angular deformity due to apposition is common. Typical large morbid mononuclear reticulum cells are scattered through these cells and typical Hodgkin giant cells are present in quantity, they tend to occur in ill defined foci. Eosinophils and neutrophils are very rare. Some iron pigment is notable.

Diagnosis. Large cell lymphosarcoma with some features of Hodgkin's disease, or Atypical Hodgkin's disease, "Lymphoblastic".

History. The patient presented with a swelling in the left parotid gland region which was provisionally diagnosed as a salivary adenoma.

The specimen is a small lymph node in the depths of which an intrusion of connective tissue containing salivary gland duct and gland acini is present. This node was reported as shewing the appearances of early Hodgkin's disease but I find myself unable to agree.

The lymph node is small, 14 X 7 X 5 mm. There is considerable persistence of the normal structure but it is masked by the hyperplasia which is present. The follicles can still be made out, they are large and some are diffuse at the periphery. Sinus tissue is well preserved in many parts. The capsule is not thickened. One trabecula bearing salivary gland elements is seen. There is some fibrosis, chiefly paratrabeular, in the form of diffuse fibrils. There is a reticulum cell hyperplasia, mainly in the pale centres of the follicles but also in the pulp. Most of the cell nuclei are small, although a few merit the term morbid mononuclear type. Rare binucleate forms are seen but no true Hodgkin giant cells. There is not any eosinophilia. Plasma cells are frequent and a few scattered neutrophils are observed. Mitoses are numerous locally - 4 or 5 per H.P. field.


Note. This patient is alive and well (November 1953). X-ray therapy has been given in several occasions. In the most recent survey available small nodes are reported to be palpable in the axillae; the spleen is not enlarged.

History. This patient complained of an axillary tumour four years ago. She was treated in Prof. J.A.G. Burton's wards and underwent an operation at this time. Thorough search of the records in the pathology department has failed to discover any reference to examination of a specimen from the patient. About two weeks previous to the present specimen a biopsy was performed but the material was so extensively necrotic that diagnosis was impossible (3451:50). The present specimen is a grossly enlarged axillary node.

This tumour is regarded as either Hodgkin's disease of highly cellular tumour-like type or a fibrifying reticulum cell sarcoma in which eosinophilia is present. A feature of unusual interest is the inclusion in the tumour tissue of a small physiologically reactive lymph node. This node is being invaded by the neoplasm at one pole. The capsule of the neoplastic node is of normal thickness but hyaline. Concentric re-formation attempts are evident but in many places invasion has occurred. Thick trabeculae are seen. Fibrosis is present, it is relatively uniform. Perivascular lamellae are common, and the fibrils present are compacted to form a coarse even mesh, the lacunae of which are filled with a delicate filigree of fibrillae. Several coralline deposits are present. The type-cell is a morbid mononuclear reticulum cell which corresponds with those of Hodgkin's disease. Small Hodgkin giant cells are common. Eosinophils are as numerous as in typical Hodgkin's disease. Mitoses are very frequent as many as 20 being present in some H.P. fields. Necrosis is chiefly individual cell type.

/Diagnosis.
87a.

Diagnosis. Atypical Hodgkin's disease, highly cellular tumour-like type, or reticulum cell sarcoma with eosinophilia. Intimately related to normal (reactive) lymph node.

History. Vague; for some time patient has noted the appearance of hard bony lumps on the long bones. His present complaint is of a painful swelling on the front of the left leg. This was diagnosed as a bone cyst and material curetted from it was sent for examination.

This is an indeterminate tumour which has affinities to reticulum cell sarcoma, myeloma of plasma cell type and Ewing's endothelioma of bone. The tumour appears very homogeneous under low power examination but high power scrutiny dispels this impression. The type-cell is superficially like a large plasma cell but the nucleus is less chromatic and oxyphile nucleoli are present. A fine reticulin mesh is present round many individual cells and groups of cells. Actual collagen is negligible. In addition typical morbid mononuclear reticulum cells and rare binucleate hyperchromatic cells are seen. Mitoses are few.

Diagnosis. Reticulum cell sarcoma, nearest to myeloma of atypical plasma cell variety.

Note. Further examination revealed 'cysts' in both femora, tibiae and left ulna but none in the skull. Bence-Jones protein was reported and then denied. Patient was traced till mid-December 1950 when he was very ill.

History. Patient has had periods of fever over the last three months latterly associated with enlargement of the right cervical lymph nodes. This was attributed to a decayed tooth but leukopenia (4,500 per c.mm.) prompted the physician to revise the diagnosis.

This specimen is a cervical lymph node shewing an exceptionally good example of classical established Hodgkin's disease.

Several lymphoid follicles at the periphery of the node survive, they have pale centres. The capsule is markedly thickened and hyaline all round the node. Concentric re-formation is well shewn and at one level a portion of the new tissue is seen in it. The exodic growth phenomenon is still discernible. Trabeculae are poorly developed. The blood vascularity is high and well seen because of congestion. The fibrosis is pathognomonon, all varieties are seen. The reticulum cell hyperplasia is characteristically protean and irregular. Large numbers of morbid mononuclear types are present together with numerous typical Hodgkin giant cells. Eosinophilia is present and many plasma cells can be found. Neutrophils are discernible throughout the node and are more concentrated round small focal necroses. Mitotic activity averages 2 per H.P. field.

Diagnosis. Hodgkin's disease, Typical. Whole node not implicated.

History. The patient presented with a painless swelling in the left groin which had developed over the last few months.

This is a fibrifying reticulum cell sarcoma with pleiomorphic cells which has possibly arisen from a follicular lymphoma. Parts (microscopically) recall Hodgkin's disease. A further point of interest is the incomplete involvement of the node in the tumour process.

One pole of the node consists of physiologically reactive lymphoid tissue and a narrow rind of similar tissue is seen in places elsewhere, the exodisc growth phenomenon being well shewn. The tumour has an irregular follicular structure the false follicles being imperfectly separated by swathes of fibrous tissue. The capsule is thickened and some concentric re-formation is evident. Fibrosis is expressed in all forms in marked degree. The cells of the tumour are a mixture of small pleiomorphic hyperchromic morbid reticulum cells and cells shewing lymphoid features. Ill-formed giant cells are noted occasionally. Eosinophils are absent, even with Dominici's stain. Plasma cells and polymorphonuclear leucocytes are scanty. Mitoses are numerous but necrosis is limited to occasional individual cells.


History. Complaint of swellings on both sides of the neck which have been present for six months. Examination revealed enlarged, discrete, firm mobile lymph nodes which were painless.

This is an example of moderately early typical Hodgkin's disease. The characteristic node of growth is well shewn. In general the morbid cells of the lesion are just appreciably smaller than usual.

Several lymphoid follicles persist even in the anatomical medulla; several shew pale centres with normal cells. A peripheral mantle of hyperplastic lymphoid tissue is seen in one node, the exodic growth phenomenon is well shewn. The capsule is thickened but this is mainly restricted to the base of the trabeculae. The new tissue in the node is arranged in coarse irregular follicles mutually isolated by collagen deposits. Fibrosis is not yet marked but all varieties are encountered. The reticulum cell proliferation is productive of many small cells although the characteristic ones are present. The Hodgkin giant cells are small and often occur in small groups. Eosinophils are present but require special stain to demonstrate them well. Plasma cells and neutrophils are noted in small numbers. Mitoses are very few though many equivocal forms exist. Focal necrosis is seen but the areas are very small.

'Plump cellular stage, eos good'. Ref. unknown. Specimen received from Dr. W. Shaw Dunn. It has been restained. History unknown beyond the fact that the patient was a boy and is now dead.

This is an example of typical Hodgkin's disease in which the complex of the lesion is dissociated, revealing the components in simpler fashion than usual. The capsule shows irregular hyaline thickening which is marked in places, and the trabeculae are also much fibrosed. The exodic growth phenomenon is seen in one segment, where a rind of physiological lymphoid tissue persists. Concentric re-formation is not seen. Fibrosis of all types except coralline deposit is seen. Small short lengths of fibrinoid are also present, these may represent collagenous degeneration. The reticulum cell proliferation is pronounced and several different varieties are being produced. Morbid mononuclear cells are present in abundance. Epithelioid cells of large size with smaller nuclei are also seen, these have rather angular acidophile cytoplasm in which minute fibrillae can be detected. These cells tend to occur in islets and are continuous with sinus littoral cells in some cases. Typical Hodgkin giant cells are present in the usual numbers. Eosinophils are very copious, nearly all are binucleate, and conform to the blood type. Plasma cells are scanty. Polymorphonuclear leucocytes are copious, they form definite micro-abscesses, and are related to necrotic foci. Mitoses are numerous, 2-3 per H.P. field.


History. Complaint of painless lumpy swelling in the left side of the neck of unknown duration. This patient was traced till January 1951. Since diagnosis he has had Murchison fever episodes approximately every three months. Blood transfusion has been given latterly with apparent benefit.

This is an example of Hodgkin's sarcoma of the variety Warthin (1931) terms type 2 or 'reticulo-endothelioblastoma'. The specimen consists of four nodes. In some there is remarkable survival of cortical lymphoid follicles in certain areas. The capsules shew very irregular fibrous thickening, in places this is intense, in others attenuation is seen. Concentric re-formation is appreciable. Fibrosis of all types is seen, the perivascular component is constant and the fibrillar deposit is chiefly seen in 'Y'-shaped strands. The reticulum cell increase is pronounced and productive of large ill-demarcated rounded areas of morbid mononuclear and Hodgkin giant cells. In addition some epithelioid cell types are being formed. Eosinophils are very scantly. Mitoses are few, below 1 per H.P. field.

Diagnosis. Hodgkin's sarcoma (Warthin's type II). Hodgkin giant cells and allied cells proliferating to the exclusion of others, regression of eosinophilia.

History. Slightly enlarged painless lymph nodes have been present for the last 18 months in the right inguinal region. Small nodes are palpable in the left groin and also in the left axilla. This specimen consisting of inguinal nodes was originally reported as an 'early low grade type of Hodgkin's disease'.

These nodes are not sufficiently enlarged to be counted abnormal in size for a man of this age. The capsules are irregularly thickened and hyaline. There is almost universal retention of normal architecture in all three, two can only be regarded as physiologically active. The third, the largest shows several diffuse areas of pulp hyperplasia. Here fibrosis is seen round several vessels and as swathes of diffuse fibrils but the connective tissue is continuous with the hilar intrusion. Reticulum cell hyperplasia is present in the pale centres and pulp, in the latter situation pathological forms are seen including typical morbid mononuclears. Eosinophils are more numerous than in normal (non-alimentary) nodes, but the increase is only convincing with sections stained by Dominici's method. Plasma cells and neutrophils are few. Mitoses are notable in the pale centres and hyperplastic pulp.

Diagnosis. Intense reactive hyperplasia? Early neoplastic change? Regard as indeterminate.

Note. This patient did not receive irradiation, the Paul-Bunnell reaction was negative. Three years later he is alive and in good health with complete regression of glandular swelling.

History. The patient attended surgical dispensary complaining of a small firm nodule which had been present two or three months on the left temple. This was diagnosed clinically as a sebaceous cyst and was resected.

The specimen is a lymph node. Superficially the appearances suggest metastatic carcinoma simplex but prolonged scrutiny does not support this diagnosis. The capsule is much thickened and pre-existing and new trabeculae cut the node into a mosaic. The tissue in the mesh consists in most cases of a curious central aggregate of pale staining epithelioid cells which are morphologically morbid mononuclear reticulum cells. (It is my experience however that metastatic malignant melanoma and occasional carcinomata can produce cells which very closely mimic these cells).

Mitoses are present, but they are definitely less than might reasonably be expected if this is cancer, multinucleate forms are present but are few. Surrounding these pale cell masses are zones of highly reactive lymphoid tissue. Fibrosis is present in other forms besides the mesh. Coralline deposits, perivascular lamellae and diffuse fibrillae are all identified. In some of the pale cell areas argyrophile fibrillae are seen. Eosinophils are present in the lymphoid tissue in numbers well above normal. Focal necrosis is seen in some of the pale cell areas.

Diagnosis. Indeterminate. It may conceivably be reactive. It may represent Ewing's trabecular type of lymph node endothelioma. It may be metastatic carcinoma or melanoma. It may be a dissociated type of Hodgkin's disease.
Note. A primary tumour has not been discovered to date, November 1954. Patient has reported at regular intervals and has always appeared well.
Alan Wilson. Aet 23 years. A discharged soldier. Ref. G.R.I. W.S. 439551. History. This patient came under observation on account of a large swelling in his left armpit of several months duration. During his sojourn in the medical wards he was also found to have an osteoporosis of the skull and the Wassermann reaction was strongly positive.

This is Hodgkin's disease of cellular type. The affected nodes are very large, 5 X 4 X 3 cm. etc. In the mass a whole node has been infarcted. Survival follicles are seen in most nodes. The exodid growth phenomenon is very well displayed in several of the sections, but concentric re-formation of the capsule is absent; the advancing edge of the new tissue is bordered directly by reactive lymphoid tissue. The capsules are thickened and hyaline, trabecular thickening is also marked. Fibrosis of all varieties is confirmed but the process is less conspicuous than usual. The reticulum cell proliferation is quite obvious and productive of morbid mononuclear types and typical Hodgkin giant cells. In addition small islets of macrophage type cells are seen. Eosinophils are numerous, many have single spheroidal lymphocytoid nuclei. Plasma cells are present, they tend to be in small focal collections rather than diffusely scattered. Mitoses are below average in number.

Diagnosis. Hodgkin's disease, fairly cellular type, accept as typical. Faint follicular pattern discernible. Incomplete involvement of nodes.

History: Since return from South Africa in March 1951 patient has exhibited the Murchison phenomenon. This was attributed to malaria presumed to have been transmitted in Liberia at a port of call. Response to treatment was unsatisfactory. Enlarged nodes were found in the neck. The Paul-Bunnell reaction was negative and two specimens of blood taken for culture were sterile.

This is early Hodgkin's disease in which there is incomplete involvement of the nodes and striking increase of reticulum cells with corresponding reduction of other cells. The Hodgkin tissue is focal in two parts of one node, a central zone of normal lymphoid tissue separating them. The exodic growth phenomenon is apparent. The capsule is wider than normal but the collagen lamellae are still separate and not impacted. Concentric re-formation is not a feature. Trabeculae are hardly represented. Fibrosis is incipient; perivascular lamellation is inconspicuous, diffuse fibrils are scanty, and fibrillae are the chief manifestation. Coralline deposits are not observed. Reticulum cell hyperplasia is marked, it is chiefly productive of a small morbid mononuclear lymphocytoid type of reticulum cell but typical Hodgkin cells are easily found among them. Giant cells are common. The reduction of lymphocytes is pronounced and those present are immature, with big nuclei. Eosinophils are present but not numerous. Plasma cells are frequent, particularly in the marginal zone. Neutrophils are scattered through the new tissue and some are collected in small foci. Mitoses average 3-4 per H.P. field. (Vascularity of the nodes is high, traumatic haemorrhage is seen).

/ Diagnosis. 
Diagnosis. Atypical Hodgkin's disease, tumour-like. Fibrosis slight and subtle. Undue reticulum cell proliferation. Affinity to reticulum cell sarcoma.

The lesion present is typical Hodgkin's disease. The specimen consists of five different nodes, the process is at much the same stage in each. Considering the blocks are 43 years old preservation is very good. The eosinophils are often disrupted but are quite recognisable.

The capsules show pronounced hyaline fibrosis, at some points very small islets of the new tissue are incorporated in them. There is a well marked exodic growth phenomenon in three of the nodes and surviving (peripheral) lymphoid tissue is discernible. Fibrosis is pronounced; the change is shown in all the standard forms, perivascular lamellae are particularly well shown. The reticulum cell proliferation is productive of morbid mononuclear and Hodgkin giant cells. These cells tend to be aggregated in masses. The eosinophilia is characteristic. (The imperfect preservation of these cells is already noted). Plasma cells and neutrophils are numerous, the latter are more numerous in the several necrotic areas which are present. Mitoses are difficult to identify with certainty due to imperfect preservation. Note — several giant-cells of Langhan's type are present.


The specimen is a cervical lymph node, it has been examined at four levels. There is not any survival of the original lymphoid tissue. The capsule is grossly thickened and hyalinised, concentric re-formation is well illustrated. The trabeculae, which are probably new formations, are very stout but do not penetrate the node regularly nor deeply. Fibrosis is well advanced, all forms are displayed and the perivascular component is the most obvious. The reticulum cell proliferation is of the usual order with the production of typical morbid mononuclear and Hodgkin giant cells. Eosinophils are copious and many lymphocytoid types can be found. Plasma cells are identified but are scanty. Neutrophils are numerous throughout the cellular part of the lesion, many are pyknotic especially those in an infarct which is present. Mitoses are of the usual order 1–2 per H.P. field.

Diagnosis. Hodgkin's disease, Typical. Late, sclerosing markedly.

History. Patient developed an intractable pruritus. Examination revealed enlarged lymph nodes in the left posterior triangle of the neck.

This is an example of tumour-like Hodgkin's disease. The cellular pleiomorphism is less arresting than usual and the typical capricious heterogeneity is lacking. In the main small cells predominate. The exodisc growth phenomenon is well shewn, a rind of physiologically reactive lymphoid tissue is seen at the periphery and concentric reformation of the capsule is seen at points. The true capsule is normal except where the new tissue actually reaches it, here thickening is present. Invasion is confirmed at two points. Fibrosis is observed in all forms except coralline deposit, it is relatively slight. The reticulum cell proliferation is productive of very numerous small cells which morphologically are intermediate between the typical reticulum cell of Hodgkin's disease and an early lymphocyte. They are almost 'normal' reticulum cells but are excessive in number. Typical morbid forms and scanty Hodgkin giant cells are present. Eosinophils are very copious and include many lymphocytoid forms. Neutrophils are scanty and plasma cells are also difficult to find. Necrosis is limited to occasional individual cells. Mitotic activity is of the usual order.

Diagnosis. Atypical Hodgkin's disease, tumour-like type.

Homogeneous growth, reduced fibrosis, invasion of capsule. Uniform small reticulum cells.

History.  Recurrence of cervical lymph node enlargement.  Previous lymphadenopathy which was irradiated.  There is not any record of a biopsy having been taken.

This is an inguinal lymph node shewing established typical Hodgkin's disease.  The lesion does not involve the whole node; in parts it is more cellular than usual.  The cellular components of a crescentic marginal portion of the node are normal; there is intense lymphatic oedema of the sinuses in it.  The demarcation of the tumour tissue is sharp at some points but almost imperceptible at others.

The capsule is much thickened and hyalinisation is proceeding, the exodic growth phenomenon is perfectly shewn and concentric re-formation is evident.  Deposits of the new tissue are seen in the capsule but are probably isolated by the last-named process.  The trabeculae are extensively fibrous.  Fibrosis of all types except for coralline deposits is well shewn.  The reticulum cell proliferation is marked locally, large morbid mononuclear types being most numerous.  Typical Hodgkin giant-cells are numerous.  Two Langhan's giant cells are seen.  Eosinophils are copious and plasma cells are frequent too, neutrophils are chiefly limited to the necrotic foci.  Mitotic figures are present in normal numbers.

Diagnosis.  Hodgkin's disease, Regarded as typical, Slightly more cellular than usual, but only locally, incomplete involvement of node.
This is an example of a tumour-like variety of atypical Hodgkin's disease. The lesion is uniformly cellular, fibrosis is sparse and eosinophils are absent. There is not any survival of normal lymph node structure. The capsule is incomplete in the specimen but is thickened where it is preserved. Local concentric re-formation is detected. Fibrosis is present but is slight; most of the collagen is fibrillar and derived by symplasmic transformation. The other forms are meagre. Reticulum cell increase is general and productive of many large morbid mononuclear cells with some epithelioid and lymphoid types. Hodgkin giant-cells are present in about the usual numbers. Eosinophils are absent and plasma cells are hard to find. Neutrophils are unusually numerous, scattered diffusely, and in areas of early necrosis. Mitoses which are unequivocal are few.


This is highly cellular tumour-like Hodgkin's disease which might reasonably be regarded as malignant from the histology. It would be equally correct to term the lesion pleiomorphic reticulum cell sarcoma.

The capsule is of normal thickness but attenuated in places; at one point it is over-run by the new tissue and the latter is growing in the peri-nodal areolar tissue. Throughout the node small survival follicles of solid type are discernible, the exodic growth phenomenon is seen. Trabeculae are not present. Fibrosis is slight; perivascular lamellation is seen round several capsular vessels, fibrils and fibrillae are scanty although reticulin fibrillae are much in evidence, the coralline form is absent. Reticulum cell increase is pronounced and general. Most of the cells are small but have hyperchromatic nuclei. Typical morbid mono-nuclear cells are copious. Multinucleate giant-cells are present but the cell outlines are indistinct giving an impression of smallness. Eosinophils are present but in small numbers, plasma cells are likewise low. Necrotic polymorphs and small lymphocytes are numerous. The mitotic figures are unusually frequent 5-7 being seen in many H.P. fields. Necrosis is limited to individual cells.

Diagnosis. Atypical Hodgkin's disease, highly cellular tumour-like type. Transgression of capsule, fibrosis slight, eosinophils low.

The material available is unfortunately not in good condition, it is traumatized and badly fixed.

The lesion corresponds most nearly to follicular lymphoma but is shewing mild eosinophilia and fibrosis of Hodgkin type. The capsule is possibly slightly thickened since where it is preserved the collagen is hyaline. It is not possible to pronounce upon the presence of infiltration.

The new tissue is disposed in false follicles some of which are incompletely outlined by compacted collagen fibres. Fibrosis is expressed as above; in several areas coralline deposit is notable. The other forms are doubtfully present. The type-cell is a reticulum cell shewing definite lymphoid type nuclei, individual morbid mononuclear types are identified and several Hodgkin giant-cells can be made out. Eosinophils are present in small foci, most are of mononuclear lymphocytoid variety. Plasma cells and neutrophils are very few. Mitoses are very scanty. Necrosis is limited to individual cells.

This is moderately early established Hodgkin's disease; the remarkable feature is that the affected node is small, 6 X 5 X 5 mm, and provides an exception to the general rule that large nodes are more likely to furnish reliable diagnosis.

Inconspicuous fragments of marginal sinus tissue and peripheral lymphoid tissue persist but the remains are insufficient to illustrate the exodic growth phenomenon. The capsule is uniformly thickened being fully 0.6 mm. wide. It is hyaline and infiltration is not seen.

The new tissue is very faintly folliculoid in arrangement, a pattern just detectable with very low power (X10). Fibrosis is expressed in all forms but perivascular lamellation is the conspicuous one. The proliferated reticulum cells are highly pleiomorphic, ranging from cells identical with fibroblasts to typical Hodgkin giant-cells. Eosinophils are present in characteristic numbers, and there is a diffuse scattering of neutrophils; plasma cells are hard to find. Mitoses average under 2 per H.P. field. Incipient infarction is seen at one pole.

Diagnosis. Hodgkin's disease, typical.
Mrs. Jean McIntyre. Ref. G.R.I. W.S.(C) 46:45.

This is an example of Hodgkin's disease at a late stage. Features of interest include the persistence of eosinophilia and relative overgrowth of the morbid reticulum cells in the surviving foci of the cellular parts of the lesion.

The capsule and trabeculae are very prominent and intensely sclerosed. This collagen formation is irregularly cribiform, and the spaces are occupied by the cellular tissue. All forms of fibrosis are encountered. The reticulum cell increase is particularly striking in foci, several high power fields approximating to mixed cell sarcoma. Large morbid mononuclear types predominate but there are plenty of characteristic Hodgkin giant-cells as well. Eosinophilia is intense especially at the margins of the cellular foci. Mitoses are still present in about the usual numbers. Neutrophils are copious and tend to be focal. Necrosis is confined to individual cells.

This is typical Hodgkin's disease which is involving part of the lymph node only, a crescentic marginal portion of original structure remains. The lesion is characterised by very numerous giant cells.

The capsule shews pronounced local thickening but this change is absent opposite the crushed normal lymphoid tissue. Concentric re-formation is absent. The demarcation of the new tissue is abrupt and an artificial cleft divides it from the normal for some distance. This phenomenon emphasizes the focal character of the disease. Fibrosis is expressed in all forms, but the coralline deposits still appear feathery in places. Trabecular thickening is gross. The reticulum cell proliferation is profuse with many aberrant forms including numerous mis-shapen Hodgkin giant-cells. Eosinophils are copious, their distribution is very even throughout the lesion. Plasma cells are scanty but many neutrophils can be seen. Mitoses average 2–3 per H.P. field. Necrosis is limited to individual cells. The sharp demarcation of normal and neoplastic tissue permits easy comparison of the lymphocytes in both formations; those in the tumour are distinctly larger with slightly less deeply staining nuclei.

Diagnosis. Hodgkin's disease, typical, (large cells more numerous than usual). Partial involvement of node well shown.

History. Patient presented with painless glandular masses in both groins.

The material is poorly preserved. The capsule is widely avulsed and tears are present. This is Brill's disease which is sclerosing. All normal structure is replaced by mutually contiguous large false follicles. Some are poorly outlined by para-vascular collagen and in the remaining pulp interstices diffuse fibril formation is seen. The type cell is the usual intermediate reticulum cell/lymphocyte precursor seen in the disease. A very few typical morbid mononuclear reticulum cells are present and one (!) Hodgkin giant-cell. There is striking absence of all granular series leucocytes and also of plasma cells. Mitoses are numerous 2-3 per H.P. field. By silver impregnation definite increase of perifollicular reticulin is apparent.

Diagnosis. Brill's disease; sclerosing in places.

(i) This is an incomplete lymph node from the sub-maxillary region, small portions of salivary gland are present. There is not any persistence of normal structure. The tissue is entirely given over to reticulum cell sarcoma of clasmatocytic stem cell type. The tumour shews a very faint follicular pattern in places, the capsule where preserved is transgressed and is being destroyed. Fibrosis is visible but only as slender fibrillae. The total collagen is small.

The tumour cells are rather small morbid mononuclear cells with typical amphophilic spheroidal nucleoli in their quasi-vesicular nuclei. The nuclei are 12-16 μ. diameter. A few large types and giant-cells are to be found. Granulocytes are absent and plasma cells are very few. Mitoses are very frequent often 10 are visible in H.P. fields.

(ii) The material is similar in all respects to (i).

Diagnosis. Reticulum cell sarcoma, clasmatocytic stem cell lymphoma. Many individual cells identical with those seen in Hodgkin's disease, fibrosis slight, faint follicular pattern.

This is a cervical lymph node showing typical Hodgkin's disease, the eosinophilia is just within the limits of acceptability. A feature of interest is the uniform immaturity of the lymphocytes, direct comparison with normal small lymphocytes is afforded by the presence of survival islets. Capsular thickening is irregular but gross where it obtains. Concentric re-formation is visible. Fibrosis is most extensive, the trabeculae are large and coarse, and all other forms are encountered. The origin of coralline deposits from compacted fibrils is clearly indicated here. Reticulum cell increase is pronounced and many grotesque forms are seen, necrosis of these cells is common. Typical Hodgkin giant-cells are numerous. The eosinophilia is less than usual, neutrophils are little in evidence. Mitoses are rare — many H.P. fields are devoid of them. There is not any focal necrosis.

Diagnosis. Hodgkin's disease, typical. (Eosinophils low).

Fibrosis very characteristic.
This is uniform cellular tumour-like Hodgkin's disease. The node is incomplete due to artefact but all the tissue present is implicated. The capsule exhibits uniform thickening of moderate degree, true trabeculae are absent. Fibrosis is chiefly in the form of perivascular accretion, the other forms are tenuous and slight. The reticulum cell increase is relatively slight and the cells are dispersed through a matrix of numerous immature small lymphocytes. In addition to the distinctive morbid mononuclear and Hodgkin giant-cells there are many small aggregates of epithelioid cells, consisting in most cases of 10-15 cells. Their cytoplasm is undergoing collagenous transformation to give coralline deposits. Eosinophilia is of the usual order but the cells tend to be focal. Plasma cells and other granulocytes are scanty. Mitoses are meagre and so far as cells can be identified in this biological phase they are in small cells probably lymphocyte precursors. Necrosis is seen in occasional morbid reticulum cells.

Blood vascularity of this lesion is high.


This is Hodgkin's disease of the soft cellular variety and it is shewing certain tumour-like features. In a large thick walled lymphatic in the adjacent areolar tissue the cells of Hodgkin's disease are visible; the only missing element is true connective tissue, the cells are set in an acidophile granular fibrin-like matrix.

The capsule is intact, thickening is present but it is within the wider limits permissable in inguinal nodes. Deposits of the new tissue are seen in the hilar intrusion portion of the capsule, but they appear to be isolation phenomena rather than infiltrative. Fibrosis is slight; the only manifestation which is easily obvious is perivascular lamellation, but scrutiny reveals fine fibrillar production in the pulp. The reticulum cell proliferation is more florid than normally seen in Hodgkin's disease, the typical morbid mononuclear and Hodgkin giant-cells are present in quantity. Eosinophilia is marked, some are of the lymphocytoid type. Plasma cells are absent but scattered polymorphs are seen. Mitoses are rather above average.


History. Complaint of pain in the left loin and groin, present for several weeks. Examination revealed enlarged nodes in the left groin.

This lesion is a peculiar mixture of Hodgkin's disease and Brill's disease. Both processes are clearly recognisable and separable in different parts of the nodes.

The capsules have been widely avulsed by artefact but where present they shew hyaline thickening. Concentric re-formation is discernible but normal lymphoid tissue is absent. A fairly uniform giant false follicle pattern with the characteristic shrinkage clefts is seen in most of the tissue but some areas reflect typical Hodgkin pattern. Fibrosis of all types is seen, it is chiefly exhibited by the Hodgkin lesion; at one area deep in the node a uniform sheet of proliferating fibroblastic reticulum cells is present. Elsewhere the distinctive morbid reticulum cells of both lesions are encountered. Typical Hodgkin giant-cells are present and tend to be focal. Eosinophils are present in adequate numbers in the Hodgkin areas. The mitoses are numerous. The lymphocytes throughout are immature in type.

Diagnosis. (Hodgkin's disease, almost typical.
(Brill's disease.

Representation is about equal.
Mrs. Betty Andison. Aet 66 years. A housewife. Ref. G.R.I. W.S. 1732:45. History. This patient exhibited the Murchison phenomenon and complained of pruritus of the back. She was very ill and weak and a swelling was found in the neck.

The specimen is a cervical lymph node, shewing late typical Hodgkin's disease.

There is not any survival of original node structure. The node capsule as received is only moderately thickened, it is probably incomplete. Hyalinisation is almost total. All forms of fibrosis are pronounced, particularly coralline deposits. Reticulum cells are relatively few and many appear almost physiological. The typical varieties including Hodgkin giant-cells are nevertheless present. Eosinophils are lower than in typical young lesions but plasma cells are more numerous. Mitoses are very infrequent. The lesion as a whole has an indolent burnt-out appearance but as usual some small active-looking foci can be found.

Diagnosis. Hodgkin's disease, typical, late.

This is an example of established Hodgkin's disease which is typical but shows a few minor deviations.

The entire node is involved. The new tissue is arranged in an irregular folliculoid manner in a coarse fibrous mesh. The capsule is very greatly thickened and extensively hyaline. Fibrosis of all forms is encountered but fibrillary deposit is much less conspicuous than the grosser manifestations. The reticulum cell proliferation is florid with the production of numerous intermediate cell-types in addition to the morbid mononuclear and Hodgkin giant cells. Several atypical giant-cells closely similar to foreign body type are noted. Eosinophilia is extreme, densely packed masses of these cells are seen in many parts. Plasma cells and neutrophils are scanty. Mitoses average 1 per H.P. field. Necrosis is absent, but haemosiderin is seen in small amount in minute foci.


This is a good example of fully established typical Hodgkin's disease with extensive fibrosis and yet only involving part of the node. A crescentic portion of physiologically reactive lymphoid tissue is present at the periphery of one side. In addition a survival follicle with a pale centre is located deep in the node. The capsule is very markedly thickened and hyalinised, except where it covers normal tissue. Fibrosis in all forms is seen, the perivascular deposits are huge. The reticulum cell proliferation is characteristic and productive of typical morbid cells. Hodgkin giant-cells are numerous and tend to be grouped. Eosinophilia is pronounced, and plasma cells and neutrophils are found with ease. Mitoses average 1 per H.P. field. Necrosis is absent.

Diagnosis. Hodgkin's disease, typical. Incomplete involvement of node despite maturity of the lesion.
Robert Law. Aet 34. Ref. G.R.I. 289:46. History. Swelling in the neck. (Note: Hodgkin's disease was diagnosed clinically in 1943, biopsy was not performed).

This is Hodgkin's disease which is typical except for the presence of many focal aggregates of epithelioid cells. The entire node is diseased. The capsule is widely lost by artefact, in the surviving fragment thickening is not appreciable. Fibrosis is relatively scanty but all varieties are seen. The reticulum cell increase is notable, many small scattered islets of epithelioid cell types are being produced and collagen fibril is being produced in these foci. The characteristic morbid mononuclear and Hodgkin giant-cells are present, the latter are distinctly scanty. Eosinophils are also rather lower than usual. Plasma cells and neutrophils are rare. The mitoses average 1 per H.P. field. Focal necrosis is absent. A notable feature is the comparative immaturity of the lymphocytes in the lesion.

Diagnosis. Slightly atypical Hodgkin's disease. Many foci of epithelioid cells. Eosinophils low. The lesion has a dissociated appearance.

Note. This patient died in G.R.I. See special P.M. series.
The appearances seen in this example of Hodgkin's disease are rare. The lesion is apparently confined to the pulp tissue of the node; many lymphoid follicles, both solid and with pale centres are dispersed through the gland. The intervening tissue is a tumour-like homogeneous Hodgkin tissue which under low power seems curiously uniform. Sinus tissue is not recognisable.

Local thickening of the capsule is seen. Fibrosis is present in all forms but a coarse uniform mesh of fibrils is the chief manifestation. In places these form pseudo-trabeculae. The reticulum cell proliferation is profuse with a high proportion of morbid forms. The Hodgkin giant-cells are rather small. Eosinophilia is somewhat less than characteristic. Plasma cells are scanty and neutrophils are very rare. Mitoses are of average frequency. Focal necrosis is absent.

Diagnosis. Atypical Hodgkin's disease. Tumour-like, confined to the node pulp, remarkable survival of lymphoid follicles.
Fred Higgins. Aet 20 years. Ref. G.R.I. 96546. History. A swelling has been present in the neck for eight months, it is painless.

Note. This patient died at home 15th October 1948. X-ray and nitrogen mustard had very little effect.

This is an example of rapidly growing cellular Hodgkin's disease. Fibrosis is minimal in most areas. Although highly cellular, the capricious pleiomorphism is not lost.

The entire node is morbid. Capsular thickening is seen and concentric re-formation is visible at several places. The trabeculae are much thickened and often infiltrated with eosinophils. Elsewhere fibrosis is slight and restricted to slight perivascular lamellation, together with a delicate fibril filigree. The reticulum cell proliferation is productive of very numerous cells, a large number of which look like fibroblasts. The morbid mononuclear and Hodgkin giant-cells are typical. Eosinophilia is pronounced, but the other cells are scanty. Mitoses are much more numerous than usual, 5-10 per H.P. field, there is much local variation. Focal necrosis is absent.

This is late typical Hodgkin's disease. The node does not retain any normal structure. Capsular thickening and hyalinisation are very marked. The new tissue is irregularly folliculoid due to pseudo-trabeculae. Fibrosis of all forms is seen, fibrillar deposit is slight compared with the gross manifestation of the other forms. The reticulum cell moiety of the lesion evidences the characteristic pleiomorphism and the typical morbid cells are present. Eosinophils are remarkably numerous, many lymphocytoid forms are present. Plasma cells and neutrophils are present in small numbers. Mitoses average 1 per H.P. field. Necrosis is limited to individual cells.

Diagnosis. Hodgkin's disease, typical, late. Note - retention of marked eosinophilia.

This is Hodgkin's disease, while probably fairly early, it is present throughout the node. There are focal aggregates of epithelioid cells which are producing fibres. The capsule is of normal thickness and is hyaline. Trabeculae are poorly represented. Fibrosis is present in all forms but is relatively slight. Fibre production is present in the epithelioid cell foci which represent the majority of the proliferated reticulum cells. Large morbid mononuclear and Hodgkin giant-cells are present but are widely scattered; many of the epithelioid cells have 'owl-eye' nuclei with prominent oxyphile nucleoli. The eosinophilia is moderate, many lymphocytoid varieties are visible. Plasma cells and neutrophils are scanty. Mitoses are of the usual order. Necrosis affects occasional individual cells.

Diagnosis. Hodgkin's disease, almost typical. Production of epithelioid cells in small islets, fibrosis is seen in these foci.

History.  Painless swelling in the right side of the neck for ten months.  Note.  This patient received X-ray therapy with transient benefit and died early in 1949.

The lesion is moderately early but established Hodgkin's disease. The node is incompletely involved. There is a crescentic marginal rind of normal lymphoid tissue and the exodisic growth phenomenon is well shown. The non-neoplastic tissue is infiltrated with eosinophils in some places. The capsule appears normal, re-formation is just appreciable. The new tissue shows an irregular nodular habit due to gross thickening of the trabeculae. Fibrosis is well established, all forms are encountered but the coralline variety is minimal. The formation of collagen fibrillae and fibrils from morbid reticulum cells is very well illustrated. The reticulum cell proliferation is florid with production of the characteristic morbid mononuclear and Hodgkin giant-cells. Eosinophilia is pronounced, it is very diffuse. Plasma cells and lymphocytes are numerous, a few widely scattered neutrophils are present. Mitoses average 2 per H.P. field. Necrosis is absent.

Diagnosis. Hodgkin's disease, typical. Incomplete involvement of node.
This is an inguinal lymph node the seat of lymphosarcoma. The tumour cell-type is an immature small lymphocyte which is better differentiated than the large cell lymphosarcoma (lymphoblastic type of Robb-Smith).

An ill-defined rind of possible normal lymphoid tissue is seen marginally against the inner aspect of the capsule. The nuclear size of the lymphocytes is smaller here and the cells are compacted. The exodic growth phenomenon can be detected but the process is much less distinct than in many cases of Hodgkin's disease. The capsule is attenuated and infiltration right through it is visible at several points.

Reticulum cell nuclei are readily found on high power examination, typical morbid mononuclear cells and one typical Hodgkin giant-cell are present. The trabeculae are appreciably thickened; fibrosis is limited to several striking examples of perivascular deposit and small coralline formations. Haemosiderin is notable in some of the perivascular lamellae. Eosinophils are present but require much search to locate. Very few plasma cells and neutrophils are discovered. Mitoses are not numerous, and are difficult to identify with certainty because of frequent pyknosis and karyorhexis.

Diagnosis. Small cell lymphosarcoma. Slight Hodgkin-like fibrosis and rare Hodgkin type reticulum cells.
William McGowan. Aet 60 years. Ref. G.R.I. 1861:46. History. Patient came under the care of Professor Burton. Laparotomy was performed for suspected abdominal tumour, enlarged lymph nodes were discovered.

The specimen is a mesenteric lymph node. This is Hodgkin's disease of fibroblastic type which is now relatively inactive. The picture is one of extensive collagenous replacement of the node, the Hodgkin's tissue surviving in small nodules and clefts in the collagen. All trace of normal structure has gone. The capsule is thin but hyaline. Fibrosis is very diffuse and amorphous masses of collagen are seen as well as coralline deposits. The reticulum cell proliferation has apparently been chiefly productive of fibroblastic cells. Morbid mononuclear and Hodgkin giant-cells are sparse. Several Langhan's type giant-cells are present. Eosinophilia is absent - very few are found, but plasma cells with deeply oxyphile cytoplasm are frequent. Other granulocytes are virtually absent. Mitoses are not seen.

Diagnosis. Atypical Hodgkin's disease, mature diffuse fibroblastic type. Low eosinophils, a few Langhan's giant-cells.

History. Patient diagnosed as retro-peritoneal tumour. Node recovered at laparotomy.

This is a lymph node from the edge of the retroperitoneal tumour. The node still shews evidence of atrophic fat replacement. Survival islets of lymphocytes are present. The capsule is of normal thickness, it is invaded at several points and tumour deposit is identified in blood and lymph vessels. Fibrosis of all types can be recognised but it is distinctly slight. The tumour is composed of morbid reticulum cells among which 'owl-eye' nucleated types and Hodgkin giant-cells of small size predominate. Eosinophilia is absent. Plasma cells and neutrophils are scanty. Mitoses average 3-4 per H.P. field.

This is a cervical lymph node showing typical fairly cellular Hodgkin's disease.

The entire node is involved in the disease. The exodic growth phenomenon and concentric re-formation of the capsule are not seen. The new tissue exhibits a poorly defined folliculoid habit. New young imperfectly collagenised swathes of connective tissue are leaving the capsule at irregular intervals. Most of the fibrosis present is perivascular, diffuse fibril formation is slight; similarly fibrillae are obscure but present. Coralline deposits are not seen. The reticulum cell increase is readily apparent and many morbid forms are being produced. Morbid mononuclear types are plenteous but typical Hodgkin giant-cells are sparse. Eosinophilia is marked. Plasma cells are notable also and several giant forms are present. Neutrophils are sparse. Mitotic figures average 1 per H.P. field. Necrosis is slight and restricted to individual cells. Haemosiderin deposit is identified. The lymphocytes are decidedly immature.

Diagnosis. Hodgkin's disease, typical.

History. Patient presented with a painless swelling in the right side of the face. The initial biopsy was reported as sinus hyperplasia probably inflammatory, the second established the diagnosis of Hodgkin's disease. The patient died 6th January 1948.

Unfortunately both specimens are damaged and the material is difficult to study.

The earlier one reveals one very definite morbid feature - marked thickening of the capsule. Where preservation is reasonably good a certain loss of normal architecture is seen, together with the appearance of scanty large morbid mononuclear reticulum cells in the pulp. Otherwise non-specific change is all that can be reported. In the second specimen fibrosis is apparent as fibrils, fibrillae, and perivascular lamellae. Reticulum cell proliferation is easily confirmed and typical Hodgkin giant-cells are present. Eosinophils are few, plasma cells and, neutrophils are little in evidence, the last named are seen in the vicinity of small haemorrhage in the new tissue. The lymphocytes are large immature types. The new tissue has a homogeneous tumour-like appearance.


History.  Patient had a small painless swelling in the left side of the neck which had been increasing in size for a year.  She was treated with irradiation but responded badly and died January 1949.

This is cellular atypical Hodgkin's disease involving a cervical node. The entire node is diseased. Capsular thickening and hyalinisation are marked, there are deposits of the cells of the new tissue in lymphatics. Concentric re-formation of the capsule is seen in places. The trabeculae are much thickened but do not penetrate deeply. Fibrosis in the node substance is relatively slight. Perivascular lamellation, fibril, and fibrillary deposits are all sparse. The reticulum cell increase is florid, most approach a macrophage in appearance and some of these cells occur in small plasmodia. Large morbid mononuclear type and typical Hodgkin giant-cells are conspicuous. Eosinophils are all but absent, plasma cells and neutrophils are not diffused through the new tissue. Several massive infarct-like areas of necrosis are ringed with neutrophils giving an abscess picture. Mitoses are numerous, they average 3-4 per H.P. field.


History. Intermittent diarrhoea and constipation for 15 months. Has lost four stone in weight and is anaemic. Laparotomy revealed enlargement of the abdominal lymph nodes.

The specimen is a mesenteric node 5 mm. diameter. It is completely replaced by rather homogeneous Hodgkin's disease tissue.

The node is enclosed in a slightly thickened capsule. At one pole a crescentic area of normal follicles, sinuses, and cords persists. The exodic growth phenomenon is well seen. The new tissue is coarsely folliculoid. Fibrosis is chiefly expressed by diffuse fibril formation; perivascular and fibrillar components are slight, and the coralline form is not seen. The reticulum cell increase is pronounced with a striking reduction of small lymphocytes. Most have small hyperchromic nuclei but large morbid mononuclear cells are present. Typical Hodgkin giant-cells are conspicuous because most of them are dead. Eosinophils are rather low in number, lymphocytoid types are notable. Plasma cells are rare but polymorphs are common especially near necrotic foci, of which there are several. Mitoses average 3-4 per H.P. field.

Diagnosis. Hodgkin's disease, probably typical. Fibrosis and eosinophils low, but this is common in mesenteric nodes. Pleiomorphism less than usual.
James Lyttle. Aet 63. Ref. G.R.I. 241:47. History. A swelling in the right side of the neck for 2½ years; glandular masses found in both sides of the neck. (Traced up till 9th November 1949 when he was dyspnoeic and had a psoriasiform lesion of the skin).

This is cellular Hodgkin's disease which almost completely involves the node.

The capsule is normal where it is preserved. Immediately beneath it the marginal sinus is intact at several points, it is filled with normal looking macrophages. (This is an unusual finding; sinus tissue is lost earlier than follicular or cord components as a rule). The exodic growth phenomenon is not appreciable. Fibrosis is slight, all forms except coralline are discernible. The reticulum cell proliferation is marked and there is production of many macrophage type cells which are often aggregated into islets. The morbid mononuclear and Hodgkin giant-cells are well represented. Eosinophils are not numerous. Plasma cells and neutrophils too are meagre. Necrosis is limited to occasional individual cells. Mitotic figures average 2 per H.P. field.

131. Serial 131.


History. Sustained a blow on the left testicle about eight weeks ago, the related groin lymph nodes swelled up and then the nodes in the opposite side enlarged. Further nodes enlarged in the left side of the neck and left axilla. All remained rather soft. After diagnosis X-ray therapy was given with a poor response, after progressive deterioration he died 6th October 1947. Total duration eight months.

This is tumour-like Hodgkin's disease in a cervical lymph node. At the periphery of the node at one level examined 13 survival nodules are present, several have pale centres but the component cells of those are not entirely normal. The exodic growth phenomenon is no longer apparent. The new tissue is very uniform and diffuse without any indication of nodular arrangement. The capsule of the node is thin, in some parts it is even attenuated but actual invasion is absent. The few trabeculae present are small but appreciably thickened. Fibrosis is slight on the whole. Perivascular lamellation is seen, fibrils and fibrillae are also apparent. Coralline deposits are lacking.

The reticulum cell increase is very pronounced, many of the nuclei are of epitheloid or fibroblastic type. Numerous morbid mononuclear types and many small but otherwise typical Hodgkin giant-cells are easily found. Eosinophils are extremely numerous often presenting in small abscess-like foci. Plasma cells are present in moderate numbers. Neutrophils are distinctly scarce. Mitoses average 1-2 per H.P. field. The lymphocytes in the lesion are immature with larger paler staining nuclei than usual.

Diagnosis. Atypical Hodgkin's disease, tumour-like. Fibrosis slight, very uniform type of growth.

History. In February 1947 she felt sick and complained of pain in the left shoulder. She also drew attention to a small lump in the left supraclavicular fossa. Examination revealed much deep lymph node involvement. She received X-ray therapy but died 2½ years after diagnosis (29th September 1949).

This is a cervical lymph node showing late typical Hodgkin's disease. No vestige of normal structure remains. Capsular and trabecular fibrosis are very pronounced. The cellular part of the new tissue is distributed in follicular fashion through the interstices of the coarse fibrous mesh formed from the capsule and trabeculae. Fibrosis in all forms is met. The reticulum cell proliferation is of the usual order and the characteristic morbid forms are readily identified. Eosinophils are still numerous and are commonly concentrated in the fibrous tissue, some lymphocytoid types are noted. Plasma cells and neutrophils are easy to find, the latter are frequent round the multiple small necrotic foci present. Mitoses average 1 per H.P. field. A small amount of haemosiderin is present in the node.

Diagnosis. Hodgkin's disease, typical, late. Retention of eosinophils.
1201:47 July, 1860:47 October. History. Swelling in the neck and
axilla of several months duration. Peripheral blood normal. The first
biopsy was reported as inflammatory lymphadenitis.

The appearances are very similar in both specimens but invasion is
pronounced in the second.

The is reticulum cell sarcoma arising from a follicular lymphoma,
the original pattern is being lost. Normal lymphadenoïd structure is not
present. The capsule is of normal thickness where it is preserved but it
is widely over-run and destroyed. Fibrosis is present; new connective
tissue is discernible as swathes of fibrils, which are occasionally
compacted round many of the false follicles. In addition there is some
diffuse fibril formation and fibrillae are appearing in the non-follicular
parts of the tumour. The reticulin impregnation reinforces this inter­
pretation. The tumour is composed of intermediate reticulum/lymphocyte
cells, the majority tending to the reticulum cell nucleus prototype with
wrinkled membranes. Large mononuclear forms and small multinucleate
giant cells are present but sparse. Eosinophils can be found but are
certainly scanty. Plasma cells and neutrophils are occasional.
Mitoses are numerous on the whole but there is much topical variation,
some H.P. fields shew as many as 6. Necrosis is limited to odd
individual cells.

Diagnosis. Reticulum cell sarcoma. Arising from follicular
lymphoma, sclerosing in parts, chiefly where follicles are disappearing.
Fields can be found which mimic Hodgkin's disease.

History. For the past year has had a lump of variable size in the left axilla. Till recently had palpable glands in the neck on the same side. The present specimen is from a mass of very large nodes, total weight 186 gm, removed from the axilla.

This is Hodgkin's disease which is typical in all respects except that eosinophils are almost absent. The lesion is universal, normal structures are all lost. The new tissue is coarsely and irregularly nodular, the cellular parts resting in the spaces of the fibrous grid. Where it is preserved the capsule is grossly thickened, some fusion with adjacent nodes is perceptible. Hyalinisation is not complete, in the trabeculae, plump fibroblastic cells are still copious. Fibrosis of all types is encountered. The reticulum cell proliferation is florid, in addition to morbid mononuclear and Hodgkin giant-cells there are many cells with characters midway between epithelioid cells and fibroblasts. Eosinophils are only discernible after much search. Plasma cells are numerous, quite frequently in small islets. Scanty neutrophils are diffuse through the tissue. Mitoses average 1 per H.P. field. Necrosis of focal type is not in evidence.

Diagnosis. Atypical Hodgkin's disease. No eosinophilia, otherwise typical.

History. See also Serial 73. The present specimen consists of two nodules removed from the scalp.

There is not any evidence that these deposits are growing in lymphadenoid tissue. The tumour presents similar appearances to those observed in the previous material but sclerosis is much better developed. The affinities to Hodgkin's disease are remote but the tumour is reminiscent of cellular fibrosarcoma in many parts.

Note. This specimen should properly belong to Serial 73; the identity of the second biopsy was not realised at the time due to the patient's name having been mis-spelt.

Diagnosis. Reticulum cell sarcoma, sclerosing. Pleiomorphism of nuclei diminishing.

History. Swelling of the left groin of six weeks known duration. The spleen was enlarged and soft. Ascites was present and the patient was very ill. The specimen is a large inguinal node 5 X 4 X 2 cm.

This is an example of large cell lymphosarcoma in which fibrosis is relatively conspicuous. There is not any survival of normal structure. The tumour is growing in a curious folliculoid fashion like the overlapping lobules of cod-roe. The lobules are oval and massive, approx. 1.5 X 2.5 mm. The capsule is appreciably thickened and the fibrous augmentation is even better displayed by the trabeculae. Infiltration of the capsule by tumour cells is prominent at most parts. Fibrosis in the tumour is chiefly perivascular and in the form of fibrils; for a tumour of this nature it is well above the amount conceded by orthodox descriptions. The type-cell of the neoplasm is a reticulum cell shewing early lymphoid differentiation. The nuclei are spheroidal, moderately leptochromatic and vesicular. The cytoplasm is usually distinct but plasmodia are common in focal areas. Eosinophils, plasma cells and neutrophils are absent. Mitoses average 3-4 per H.P. field. Necrosis is not a feature.

Diagnosis. Large cell lymphosarcoma. Folliculoid pattern discernible. Sclerosing slightly.

History. Enlarged painless lymph nodes have been present in the left side of the neck for four months.

This is a cervical lymph node which I believe to be an example of Benign Hodgkin's disease (Harrison 1952). It was originally reported as early Hodgkin's disease.

The capsule and trabeculae are greatly thickened and hyaline, this constitutes the main alteration visible - it is easily seen on naked eye examination of the section. The lymphoid tissue is distorted in a subtle fashion; solid follicles and ones with pale centres are present especially peripherally, but they are not normally disposed. Centrally the pulp has overgrown and exodic growth is seen. Fine fibrosis is not present but coralline deposits are numerous. Reticulum cell proliferation is seen in the pale centres of follicles and also in the pulp but in the latter site it is inconspicuous. Scanty large mononuclear forms and two Hodgkin giant-cells are identified in one section. Eosinophils, plasma cells, and neutrophils are conspicuously absent. Mitoses are very scanty indeed. Necrosis is absent.


History. Patient has had a swelling in the left side of the neck for several months, its size has been static and it is painless. It was diagnosed clinically as a sebaceous cyst and resected.

This is early active Hodgkin's disease in a cervical lymph node. Under low power poorly demarcated follicular aggregates of lymphocytes can be seen. High power examination dispels this impression of surviving normal tissue. The lymphocytes are immature with large leptochromatic nuclei. Set in among them in multiple irregular islets are areas of cells simulating pale centre tissue. Some of these reticulum cells have large morbid nuclei. The exodic growth phenomenon is not apparent and concentric re-formation of the capsule is absent. The capsule is of normal thickness but hyalinised. Fibrosis is seen in increased stoutness of the trabeculae and as fibrils, fibrillae and coralline deposits. Perivascular formation of collagen is lacking. (There is an absence of sizeable vessels). The reticulum cell proliferation outwith the folliculoid formation noted above is marked. Scanty epithelioid cells are seen, but numerous morbid mononuclear types and Hodgkin giant-cells are identified. The latter are small but otherwise typical. Eosinophils are numerous and are diffuse. Plasma cells are absent - an unusual event. Polymorphs are likewise scanty. Mitoses average 2 per H.P. field. Necrosis is absent.

Diagnosis. Hodgkin's disease, typical, early, active. ? early disorganisation of follicle tissue of node. Primary origin in node pulp is likely. Affinities to reticulum cell sarcoma.

It is difficult to decide which is the most appropriate category for this example of lymph node tumour. The capsule of the node is incompletely preserved but the remnants are thin. Trabeculae are hardly represented. Portions of marginal sinus and several peripheral follicles are identified; in addition fragments of trajectorial sinus tissue survive. The bulk of the node however is devoid of normal architecture and composed predominantly of small cells. This diminution in size is particularly noticeable in the small lymphocytes which conform to the type seen characteristically in some lymphosarcomata. Fibrosis i.e. actual collagen formation, is absent, but reticulin is definitely increased as a fine pervading filigree. Scattered through the small lymphocyte population many small morbid mononuclear and multinucleate giant-cells are discerned. Poorly demarcated paler staining areas visible low power are seen to be made up of small angular cells with ground-glass oxyphile cytoplasm and lymphocytoid nuclei. Eosinophils are almost absent, plasma cells are scanty. Neutrophils are doubtfully present. Mitoses are very infrequent most examples turn out to be pyknotic fragments.


This is tumour-like Hodgkin's disease in an axillary lymph node. The node is totally diseased. The capsule shows thickening and islets of new tissue are present in it. Trabeculae and swathes of new connective tissue cut the cellular components into irregular nodules. In the nodules the lesion is curiously regular and uniform with indistinct whorled arrangement. Fibrosis is moderately early in maturity and present in all forms except coralline deposit. Reticulum cell proliferation is universal and mostly productive of a small fibroblastic type of cell. The characteristic morbid mononuclear cells and Hodgkin giant-cells are present in rather small numbers. Eosinophils are scanty but neutrophils, often as micro-abscesses are very numerous. Plasma cells are scanty, most are located at the periphery of the node. Mitoses are few, below 1 per H.P. field. Many infarct-like areas of necrosis are notable both peripheral and central.

Mrs. Agnes Tannahill. Aet 47 years. A housewife. Ref. G.R.I. 729:46, 2611:48. History. Swelling of the lymph nodes in the neck, axillae, and groins of several months duration. The initial specimen is a cervical lymph node, the second a node recovered from the vicinity of the sac of a femoral hernia which had been present seven or eight years.

This is a good illustration of the instability of follicular lymphoma (Brill's disease).

The first specimen shews considerable fibrous thickening of the node capsule. The tumour shews typical false follicles which are better seen peripherally, the pattern is indistinct centrally. There is considerable pleiomorphism of the reticulum cells both in the false follicles and central pulp. Large morbid mononuclear and Hodgkin giant-cells are readily identified, but integral fibrosis is absent and there is no eosinophilia.

In the second specimen there is almost total loss of follicular pattern. Fibrosis of all varieties is present, fibrils being the chief form. The capsule is widely destroyed and over-run by the tumour cells. The cellular pleiomorphism is enhanced a little and the cell nuclei are larger, there are proportionately more lymphocyte series cells of primitive type. Hodgkin giant-cells are present in very small numbers and are distinctly small. Eosinophils are absent, plasma cells are very meagre. Mitoses average 4-5 per H.P. field. Necrosis is not a feature. Tumour deposits are confirmed in the lumen of an artery and permeation of the wall is present.

Diagnosis. Pleiomorphic reticulum cell sarcoma. Arising from Brill's disease, sclerosing.

History. Painless slowly growing swelling in the left side of the neck, present for the past two years. Diagnosed provisionally as tuberculous lymphadenitis.

The specimen consists of several pieces of cervical lymph node which are fused. All are neoplastic except one, the normal node retains its architecture and cell structure and is almost entirely encircled by tumour. This latter phenomenon is of much interest as it supports the view that systematisation is not necessarily a feature of tumours of the reticulo-endothelial system.

The neoplastic nodes shew reticulum cell sarcoma of undifferentiated type presenting certain morphological affinities to Hodgkin's disease.

The capsules where shewn are thick and hyaline but are extensively transgressed at many points. Concentric re-formation is notable. A faint nodular habit of growth is exhibited by the tumour. Fibrosis of all forms is seen but it is moderate. The tumour cells are pleiomorphic morbid reticulum cells shewing little differentiation, free cells are present but most form imperfect plasmodia. Large morbid mononuclear types are common but Hodgkin giant-cells are scanty. Eosinophils are absent and plasma cells are very meagre. Neutrophils are hard to identify due to pyknosis. Mitoses are very numerous 8-10 per H.P. field.

Diagnosis. Reticulum cell sarcoma, sclerosing in places. Affinities to Hodgkin's disease close. A physiologically reactive lymph node is enclosed in the tumour.

The specimen is a cervical lymph node shewing atypical Hodgkin's disease in which small cells predominate and eosinophils are absent.

The node is entirely morbid. The habit of growth of the tumour is faintly nodular. The capsule is very markedly thickened and hyaline, concentric re-formation is present, and at several points there is enclosure of islets of the new tissue in it. In these latter sites the tumour does not appear aggressive. Trabeculae, which are probably new formations partially separate the nodules of the new tissue. Fibrosis of all forms is encountered but are slight as compared with the overwhelming sclerosis of the capsule. The reticulum cell proliferation is productive of very numerous small cells but scrutiny reveals large morbid mononuclear types and infrequent Hodgkin giant-cells. Eosinophils are absent. Plasma cells and neutrophils are conspicuously numerous, well above average for a pure Hodgkin lesion, it seems likely that inflammation is responsible as the necrotic foci are not very numerous nor large. Mitoses average 2 per H.P. field. The lymphocytes include many immature forms.


History. Painless swellings in the neck of three months duration.

The material is poor, the node collapsed on removal. This is a small cell lymphosarcoma. The capsule is indifferently preserved due to being over-run by tumour cells and traumatized. The exodic growth phenomenon is displayed. Fibrosis is minimal, several small scar like areas of compacted fibrils are the only manifestation of note. Nearly all the cells are definitely of the lymphocyte series and small nuclei preponderate. The few large nuclei found are blood vascular endothelial. Eosinophils are absent and plasma cells are sparse. (Haemorrhage due to trauma is seen). Mitoses are difficult to find due to pyknosis, many of the necrotic nuclei assume the characteristic lymphocyte form of bivalve type.

Diagnosis. Small cell lymphosarcoma. Affinities doubtful. (Poor specimen).

History. Referred to out-patients dispensary on account of persistent cervical adenitis. The specimen is a cervical lymph node.

This is Brill's disease in which the follicular pattern is being lost or is poorly developed. Invasive features are notable, and some fibrosis is present.

The entire node is morbid. There is subtle mimicry of the normal architecture by the pseudo-follicles which are mostly of the solid variety, and composed of cells which are closely similar to those of the pulp. Shrinkage clefts are observed in many cases. The capsule is of normal thickness, it is transgressed at several points with spread of the tumour cells into the adjacent areolar tissue. Fibrosis is just appreciable, minute representatives of all varieties are found. The type cell of the growth is an immature lymphocyte. Rare morbid mononuclear forms are to be seen but there are not any giant-cells. Eosinophils and other granulocytes are absent, plasma cells are very scanty. Mitoses average less than 1 per H.P. field. Necrosis is not a feature.

Diagnosis. Brill's disease, regard as malignant. Follicular pattern less distinct than usual. Fibrosis slight but definite.

This is an example of lymphoid tissue sarcoma in which the appearances are not appropriate to any of the common categories, the microscopical appearances did not alter very much in the observed course.

The first specimen is a small node 10 X 8 X 6 mm. There is a rind of survival tissue beneath the capsule which is fairly sharply distinguishable from a homogeneous tumour-like tissue occupying the centre. The exodic growth phenomenon is very well illustrated. In this node the capsule is normal in thickness and is not invaded. In the central new tissue fibrosis is present. Perivascular lamellae, fibrils in swathes, and fibrillae are seen. There is reticulum cell proliferation with the production of a uniform mixture of small morbid mononuclear cells and immature lymphocytes. Eosinophils are present in small numbers. Rare small giant-cells are visible. Mitoses are few.

In the second specimen, which is several centimetres across, the capsule is thicker and infiltrated by tumour, normal lymphoid relics have disappeared. The growth is similar to the previous stage except that growth is much quicker. In most areas the lymphoid propensities are greater but in a few the appearances are very close to cellular Hodgkin's disease. Necrosis is not evident.

/Diagnosis.
Diagnosis. 1. Indeterminate, ? intermediate lymphoid tissue sarcoma.

2. Indeterminate. Lymphoid tissue sarcoma (malignant) probably nearest to lymphosarcoma but has close affinity to cellular Hodgkin's disease.

History. Vague abdominal pain of three months duration. Palpable mass present in the epigastrium; provisionally diagnosed as mesenteric sarcoma. The patient died of lymphosarcoma 30th January 1950.

The first specimen is a mesenteric lymph node, the second is from the axilla. This is lymphosarcoma in which a false follicle pattern is visible in the earlier biopsy but is much less distinct in the later specimen. There is not any persistence of normal structure in either node. The capsules are thin, attenuated in places, and over-run by small lymphocytes. Fibrosis in the tumour is minimal but is slightly better seen in the later node, it is in the form of diffuse fibrillae. The type cell is an immature lymphocyte with a large moderately deep-staining stippled nucleus, however many morbid mononuclear types are visible intermingled in the second specimen. Several small multinucleate giant-cells are also present in 2. Eosinophils are absent and the other cells, plasma cells etc. are rare. Mitoses average 4-6 per H.P. field. Necrosis of individual cells is noted.

Diagnosis. Large cell lymphosarcoma. Possibly arising from follicular lymphoma. Apparently becoming more primitive (rise in proportion of reticulum cell types).

This is a perplexing case of lymph node enlargement of indeterminate character. The specimen is a large lymph node (2.5 X 2 X 1.5 cm.) from which the capsule has been widely avulsed. The original diagnosis of Hodgkin's disease is unquestionably erroneous. The appearances are more reflective of inflammatory reaction and hyperplasia than tumour but such gross enlargement is anomalous. The patient is alive and well (4½ years later). Only a small fragment of capsule persists. The node is beset with indiscriminately scattered hyperplastic germinal follicles. In many the outline of the pale centre tissue is distorted but does not shew any tendency to separate from the peripheral lymphocytes. The pulp shews irregular local over-growth which drives the follicular tissue aside in places. Several cases of marked perivascular fibrosis, and small irregular scar-like fibrous areas are also visible in the pulp. Several micro-abscesses are present most in the pulp but some apparently in follicles.

In the follicular tissue the pale centres are crowded with rapidly proliferating reticulum cells and tingible Korper are copious. The cells are chiefly physiological in size but some are morbidly large with rare binaucleate forms. In the pulp similar reticulum cell overgrowth is seen, here morbid forms are commoner but tend to be smaller than typical examples. True Hodgkin giant-cells are not identified. Eosinophils are very scanty but plasma cells abound. Neutrophils are present in the suppurative foci, pyknosis is common.

/Diagnosis.
Diagnosis. Indeterminate, regarded as probably inflammatory. Mimics Hodgkin's disease, but insufficient pleiomorphism, absence of diffuse fibril collagen, and retention of normal structure preclude acceptance of original diagnosis.

History. A painless swelling has been present in the left axilla for the past six years. This was provisionally diagnosed as a cyst and was resected, it measured 4.5 cm. diameter. The peripheral blood is normal.

The specimen is a greatly enlarged lymph node shewing lymphosarcoma with unusual features. An ill-defined rind of reactive physiological lymphoid tissue without follicles is present at the subcapsular periphery. The exodic growth phenomenon is well shewn. Deep to this reactive tissue is a uniform lymphocytic tumour which does not shew any follicular pattern. The node capsule is moderately thickened but is infiltrated with tumour cells at many points. Fibrosis is present in unusual amount for growths of this kind. Perivascular lamellation is marked, fibrils and fibrillae are also notable. Small coralline deposits are seen round several of the larger vessels. The type cell of the tumour is a small lymphocyte which is almost mature. Large morbid mononuclear reticulum cells are scattered sparsely but evenly through the growth and small islets of epithelioid cells are also found. Several typical Hodgkin giant-cells are present. Eosinophils and neutrophils are absent but a few plasma cells are visible. Mitoses average 2 per H.P. field. Necrosis is not seen.


History. Has suffered from malaise and lassitude for the last four months. Swellings have been present in the neck. The blood shews chronic lymphatic leukaemia.

This is small cell lymphosarcoma in a cervical node. The blood vessels and lymphatics of the adjacent areolar tissue are packed with small lymphocytes. The only surviving normal structure is part of the marginal sinus. (This is an unusual finding). The node capsule is thick and hyaline; infiltration is equivocal. Fibrosis is very slight, a few collagen fibrils are demonstrable round the vessels by special method. There is not any increase above normal. The type-cell of the tumour is an immature small lymphocyte, the nucleus of which is fainter staining than mature small lymphocytes. Rare morbid mononuclear cells are identified. Eosinophils and neutrophils are absent, a few plasma cells can be found. Mitoses average 2-3 per H.P. field, necrosis is absent.

Diagnosis. Small cell lymphosarcoma, almost pure.

History. Generalised lymph node enlargement present for eighteen months. Examination of the blood revealed chronic lymphatic leukaemia.

The specimen is a small cervical node 9 X 5 X 5 mm. shewing lymphosarcoma in which a very indistinct but definite follicular pattern is visible. Very little fibrosis is present.

Normal lymphoid tissue does not survive. The new tissue shews a faint micro-follicular pattern in the deeper part of the node. The capsule is of normal thickness and hyaline, infiltration is confirmed at one point but is not extensive. Fibrosis is very slight, the deposits are almost certainly hyalinised vessels.

The type-cell is an immature lymphocyte, intermediate in size between small and large lymphocyte. Rare morbid reticulum cells are present, they are widely scattered. Eosinophils, neutrophils and plasma cells are absent. Mitoses are very few, 1 per 3 H.P. fields. Occasional individual cells are necrotic.

Diagnosis. Lymphosarcoma, pure, with leukaemia. Very faint follicular pattern.

This node is inflamed and highly reactive, it also represents lymphoid tissue at the zenith of its natural activity. It is possibly an example of the poorly characterised mesenteric lymphadenitis.

The chief justifications for its inclusion in the series are the nuclear pleiomorphism of the reticulum cell proliferation in the pulp and the formation of new lymphoid tissue in and outwith the capsule. Scattered eosinophils are present, a constant finding in mesenteric lymph nodes. Plasma cells are more numerous than normal.

Diagnosis. Physiological reaction and mild non-specific chronic inflammation.

History. Generalised lymph node enlargement present for six weeks.

This is atypical Hodgkin's disease of a peculiar type. The bulk of the new tissue is composed of reticulum cells of epithelioid or fibroblastic type, and fibrosis is chiefly arising by direct diffuse symplasmic transformation. Small areas of typical Hodgkin's disease morphology are identified. Overall the picture is tumour-like. Survival follicles of lymphoid tissue, most with pale centre tissue are present widely scattered through the new tissue; they are not limited to the periphery, this supports the view that the pulp may be primarily involved. The capsule is imperfectly preserved but the pieces remaining are thickened, infiltration with the new tissue is present. Trabeculae are not present. Fibrosis is chiefly fibrillar and appears as reticulin at the margins of the epithelioid cells, much is visible with van Gieson's stain and is apparent also in the routine section. The reticulum cell proliferation is very conspicuous, most of the cells are as noted above. In addition typical morbid mononuclear and Hodgkin giant-cells are numerous. Several Langhan's type giant-cells are found, they are quite indiscriminately scattered. Eosinophils are copious and plasma cells are easily found; these features are probably enhanced by the simplification of the tissue and sparse small lymphocytes. Neutrophils are also fairly numerous and are most concentrated round minute necrotic foci. Mitoses average 1 per H.P. field.


History. Swelling in the right side of the neck which has been present for one year. The patient is a diabetic and is cachectic despite treatment.

The specimen is a cervical lymph node shewing Hodgkin's disease without eosinophils and some tumour-like homogeneity.

There is not any survival of lymphoid tissue. The new tissue shews faintly the expanding nodule habit of growth. Deposits of the tumour are visible in the part of the capsule which is preserved, they appear isolated rather than infiltrative. Thickening of the capsule and trabeculae is pronounced. Fibrosis of all types is encountered and coralline deposit is unusually well evidenced. An interesting anomaly is encountered in the fibrillary manifestation; the fibrillae take the cosin and silver well but stain very faintly with the fuchsin of van Gieson's stain. The reticulum cell proliferation is relatively uniform and the numerous large morbid mononuclear cells are evenly scattered. Typical Hodgkin giant-cells are present in the usual numbers. Eosinophils are very few and other granulocytes are sparse. Plasma cells are scanty. Mitoses are not above 1 per H.P. field. Necrosis affects individual cells, chiefly those with large nuclei.

Diagnosis. Atypical Hodgkin's disease. No eosinophils, tending towards tumour-like type.

History. A cancer of breast was diagnosed clinically five years ago. Biopsy was not performed. Deep X-ray therapy was administered. A swelling developed in the ipsilateral axilla four weeks ago. The breast and axillary nodes were resected.

This specimen was examined very thoroughly for evidence of tumour, some 20 blocks being taken and all proved negative. The node examined shews a picture which simulates Hodgkin's disease in some respects, sinus tissue is difficult to identify.

There is general retention of the normal architecture of the node, reactive pale centres of irregular outline are present in some of the follicles and in several places there is marked pleiomorphism of the pulp reticulum cells. The capsule is thicker than normal and the trabeculae partake of this change. Diffuse infiltration of these tissues by lymphocytes and plasma cells is notable. Fibrosis in the node tissue is practically restricted to perivascular thickening; the typical lamellation is absent and endarteritic stenosis is common. Rare diffuse fibrils are present in the vicinity of trabeculae. The proliferated reticulum cells assume bizarre forms, but in nearly all the nuclei are noticeably small. Multinucleate giant-cells are absent. Eosinophils are absent, neutrophils and plasma cells are above normal in number.

Diagnosis. Reactive and chronic inflammatory change. Vascular fibrosis, probably attributable to X-ray therapy.

This is an interesting example of reticulum cell sarcoma shewing a striking resemblance to Hodgkin's disease.

There is not any survival of normal lymphoid tissue. The capsule is grossly thickened and hyaline, but despite this it is widely infiltrated and transgressed by tumour which is spreading into the adjacent fat. The neoplasm is an alveolar sarcoma and shews the expanding nodule type of growth. Fibrosis is pronounced, all forms are present, but the coralline formation is the most obtrusive. It forms a coarse rather irregular mesh in the interstices of which lie clusters of tumour cells. The type-cell is a free cell conforming to a morbid mononuclear type except that it is rather smaller. The nucleus is more densely staining too, but there are also a fair proportion of typical morbid mononuclear reticulum cells. Small Hodgkin giant-cells are present. Eosinophils are absent and it is difficult to find any plasma cells or neutrophils. Mitoses are very numerous, up to 20 can be seen in H.P. fields. Necrosis is limited to individual cells and is inconspicuous.

Diagnosis. Reticulum cell sarcoma, nearest to "clasmatocytic stem cell lymphoma" but some lymphoid propensities present in the tumour cells. Fibrosis and areas of pleiomorphism very similar to Hodgkin's disease.
Columba Newcombe. Act 27 years. Ref. G.R.I. 3579:50. History. Enlarged shotty lymph nodes in the neck and axillae of one months duration. There is not any pyrexia, the peripheral blood is normal, and the Paul-Bunnell reaction is negative.

This is a small cervical node, 8 X 4 X 4 mm. There are several areas of interstitial haemorrhage and serous exudate in it. The general architecture remains normal in most areas with retention of lymphoid follicles including many with pale centres. In the deeper part and at some points at the periphery the pulp shews unusual cellular pleiomorphism. The capsule is appreciably thickened but still cellular, at the hilar intrusion, and many small lymphocytes are seen between the lamellae. Fibrosis is present round several large vessels, the largest is apparently the hilar capsule mantle and must be regarded as normal. The trabeculae are also rather more prominent than usual. Scanty fibrils and fibrillae are visible in parts of the pulp. There is considerable reticulum cell increase in the pulp. The cells produced are pleiomorphic but few are sufficiently large or have large enough nuclei to be considered morbid. Small groups of irregularly shaped epithelioid cells are seen at places and scattered singly. Eosinophils are present but not above the usual numbers. Neutrophils and plasma cells are rare. The lymphocytes are normal. Early necrosis is present in the cells in the exudate areas. Mitotic figures are rare except in the pale centre nuclei.

Diagnosis. Indeterminate, considered physiological.

Note. Alive and well three years later (27th October 1953).

History. Swelling in the posterior shoulder region of unknown duration. (It was originally stated that enlarged lymph nodes were not present but this was later revised). The patient responded badly to X-ray therapy and died 30th March 1952.

The material is a biopsy from the tumour of shoulder, it is badly preserved but permits diagnosis. The growth is Hodgkin's disease of tumour-like type. (This appearance is so real that it was in fact reported as reticulum cell sarcoma, however it conforms more closely to Hodgkin's disease).

The tissue is probably a group of small lymph nodes which have thickened fused capsules. Trabeculae are well developed both mature hyalinised bands and swathes of younger tissue. Fibrosis of all forms is encountered but coralline deposits are small. The reticulum cell proliferation is productive of very numerous immature small lymphocytes. These cloak the typical large morbid mononuclear and Hodgkin giant-cells which are readily found on close scrutiny. Eosinophils are few and tend to be focal. Plasma cells are very difficult to identify but this is certainly partly due to the poor preparations. Mitoses are likewise difficult to confirm.

Diagnosis. Atypical Hodgkin's disease, many immature lymphocyte series cells, few eosinophils.

Note. It was later learned that this patient had a severe spastic paraplegia attributed to a deposit in bone at T.7-8 level.

History. Generalised lymph node enlargement of several months duration.

This is an example of soft cellular Hodgkin's disease shewing tumour-like homogeneity. The eosinophils are much reduced. The node is an axillary one and is large; 20 X 30 X 17 mm. Rare follicles and several islets of loose lymphatic tissue persist at the sub-capsular periphery. The exodic growth phenomenon is passing off. The new tissue shews many small focal areas of necrosis or incipient necrosis. The capsule is thin and transgressed in several places. Concentric re-formation is just discernible at some points. Several well developed thick trabeculae are notable, including swathes of fibrils not yet compacted. Fibrosis of all varieties is seen, the greater part of it is in the form of fibrils and fibrillae. The latter are being formed by symplasmic transformation. The reticulum cell overgrowth is marked and lymphocytes are relatively scanty. Most of the cells are epitheliod or fibroblast types and their cytoplasmic borders are condensing to form collagen fibrillae. Typical morbid mononuclear and Hodgkin giant-cells are copious. Eosinophils are below the usual number but are present in most fields. Plasma cells are scanty but neutrophils are common especially in foci round the necrotic areas. Mitoses are below 1 per H.P. field. Necrosis is focal as already noted.

Robert Shaw. Aet 49 years. Ref. Law Hospital 506:51. History. Complaint of sudden acute colicky pain in the epigastrium. Examination revealed signs of acute intestinal obstruction. Laparotomy was performed and disclosed a tumour of the colon and enlarged lymph nodes in the abdomen. (The latter were not disturbed).

The tumour is a pleiomorphic reticulum cell sarcoma which appears to have arisen in the lymphoid tissue of the caecal wall. Fibrosis is seen in the neoplasm in the form of strands, some compacted into swathes; fibrils and fibrillae. The type cells are small morbid reticulum cells with hyperchromic nuclei. Large typical ones are present and small Hodgkin giant-cells are visible. Eosinophils are present but in small numbers. Plasma cells and neutrophils are absent from the deeper parts of the growth but appear on the ulcerated surface. Mitoses average 2-3 per H.P. field, necrosis is limited to the surface presenting in the lumen. The growth is invading the muscle wall but is not very aggressive.

Diagnosis. Hodgkin's sarcoma. Pleiomorphic reticulum cell sarcoma, corresponds closely with Hodgkin's disease in general features but frankly malignant. (Warthin Type II).

History. Painless swelling in the right side of the neck for twelve months. The specimen consists of a partly fused mass of large and small lymph nodes. Total size 9 X 6 X 6 cm.

This is Hodgkin's disease of soft cellular type which is becoming tumour-like.

There is not any persistence of normal organised lymphoid tissue but small lymphocytes are more numerous at the sub-capsular periphery. The capsule is slightly thickened locally at a few points, in the more tenuous reaches it is infiltrated with the new tissue. Trabeculae are absent. Fibrosis is distinctly light, all forms are encountered except coralline deposits, much is derived by symplasmic transformation. The reticulum cell proliferation is universal and chiefly productive of epithelioid and fibroblastic cells. Both are producing collagen. Typical morbid mononuclear cells and Hodgkin giant-cells are numerous throughout. Eosinophilia is focal, many fields being free of these cells. Plasma cells and neutrophils are common round small necrotic foci. Mitoses average 1-2 per H.P. field.

Diagnosis. Atypical Hodgkin's disease, soft cellular variety with tumour-like homogeneity. Low eosinophilia and tenuous fibrosis.
Isabella Aitken. Aet 40 years. Ref. Law Hospital 532:51.

History. A lump in the right side of the neck in the supraclavicular region for seven months. An X-ray photograph of the chest taken seven months ago was reported normal. The swelling is painless, the other accessible nodes appear normal.

This is Hodgkin's disease which is typical except that eosinophils are sparse, in all other features the lesion is classical. The example illustrates the principle that while eosinophilia is typical it probably is not an essential feature of the lesion.

Normal lymphoid tissue does not survive. The capsule shows marked fibrous thickening, much of it is still cellular especially on the inner aspect. There is intense sclerosis of pre-existing trabeculae and numerous swathes of new connective tissue have developed. Fibrosis in all forms is pronounced, the coralline deposits are seen to result from fibril compaction and vascular hyalinisation. The new tissue is disposed in irregular nodules. The reticulum cell proliferation is productive of numerous large morbid mononuclear cells and Hodgkin giant-cells. The latter are rarely very large. Eosinophils are sparse. Plasma cells are common but neutrophils are rare. Mitoses average 2-3 per H.P. field but the sections are thick. Necrosis is not a feature.

Note. A small node 1 mm. diameter adjacent to this large one shows recognisable Hodgkin's disease of the same type.

Diagnosis. Atypical Hodgkin's disease, low eosinophilia the only atypical feature.

This is best described as an intermediate reticulum cell sarcoma possibly emerging from follicular lymphoma. Several nodes are available for study and the appearances are different. In the smaller ones follicular lymphoma is present, the follicles are large, contiguous and beginning to lose their mutual demarcation; in the larger nodes this pattern is disappearing, particularly centrally. The larger nodes probably represent the later stage of the change and are therefore described.

Their capsules are of normal thickness, concentric re-formation is notable between the peripheral pseudo-follicles and the subjacent diffuse growth. Invasion and extension into the adjacent fat is present. The central tissue of the neoplasm shews fields which approximate to small cell lymphosarcoma, to reticulum cell sarcoma, and mixtures of the two. Large morbid mononuclear reticulum cells are found readily but Hodgkin giant-cells are absent. A few Dorothy Reed mirror-image cells are seen. Fibrosis is slight, it is chiefly of diffuse fibril and fibrillary form. The silver impregnation reveals many more pre-collagen fibrillae. Eosinophils, neutrophils, and plasma cells are inconspicuous, the foremost being very sparse. Mitoses are difficult to identify due to widespread pyknosis.

Diagnosis. Reticulum cell sarcoma, intermediate type, sclerosing slightly. Affinities to Hodgkin's disease very close in places.
James Hogg. Aet 52 years. Ref. G.R.I. 702:51. History. Painless enlargement of the lymph nodes and liver found on clinical examination of the patient who is considered by the dermatologists to be suffering from mycosis fungoides ('pre-mycotic' stage).

The specimen is a large axillary lymph node, 30 X 20 X 20 mm. The general architecture is preserved but modified. Sinus tissue survives and littoral cell increase is pronounced locally giving a solid packed appearance. Germinal follicles are present both solid types and those with pale centres, in some of the latter the central capillary is hyalinised. The pulp cords are intact but instead of being composed of small lymphocytes nearly all the cells are plasma cells. The capsule and trabeculae are thickened. Fibrosis is seen round vessels and there is some diffuse fibril formation. There is a great increase of reticulum cells in the inter-follicular pulp and haemosiderin is present in the cytoplasm of some. The great majority of these cells have nuclei of physiological size, a few conform to the morbid type, and rare binucleate forms are encountered. Eosinophils are present but are scanty. Mitoses are numerous among the proliferating reticulum cells of the pulp.

Diagnosis. Indeterminate. Reticulum cell proliferation pronounced in areas of the pulp, slight haemosiderosis. Plasma cells very numerous in capsule and lymph cords deep in the node. Eosinophils present. Regarded as primary, chiefly on account of the size.

History. This patient was operated on for the treatment of varicose veins of the leg. At the operation the groin lymph nodes appeared unduly prominent and three were resected.

The nodes are small, the largest being 10 X 9 X 8 mm. The general architecture is retained. The capsules and trabeculae are normal. Lymphoid follicles, some of the them with normal pale centres are present. Fibrosis is present round many of the arterioles of the node and multiple small coralline deposits are seen scattered indiscriminately through the node structure. There is marked proliferation of reticulum cells in ill-defined areas of the pulp, usually in the peripheral inter-follicular loose lymphatic tissue. While most of these cells are small and evenly dispersed a few conform to morbid mononuclear type. Eosinophils are distinctly more frequent than in normal nodes. Serial sections reveal that most of the coralline collagen deposits can be accounted for by hyalinisation of blood vessels.

Diagnosis. Physiological reactive change. Small areas which mimic, or rather understudy, Hodgkin's disease are discernible. The reacting cells are small.

History. A solitary swelling in the axilla present for several months, provisionally diagnosed as a cyst. Naked eye the tumour was considered to be a lipoma. Dimensions 6 x 3 cms, ovoid.

This is an anomalous lymphoid tissue tumour, it bears a superficial resemblance to small cell lymphosarcoma of low malignancy but remarkable fibrosis is present.

The growth is apparently a grossly enlarged lymph node. A few deformed solid lymphoid follicles are discernible at the periphery. Two have imperfect pale centres. The exodic growth phenomenon is present but there is not the reactive change of the peripheral surviving tissue which is usually seen. The capsule is slightly thickened and hyaline, invasion is not seen in the blocks (3) selected for microscopy, concentric re-formation is not seen. Fibrosis is seen in all forms but the most extensive is coralline deposit. It is characteristically similar to the type encountered in Hodgkin's disease. The type-cell of the growth is the small lymphocyte but immature slightly leptochromatic larger nuclei are common. Very few morbid mononuclear reticulum cells are seen. True Hodgkin giant-cells are absent. Eosinophils occur in small numbers. Plasma cells are present and neutrophils are seen in tiny pseudo-abscess aggregates. Mitoses are few.

Note. The coralline collagen deposits are traceable to vascular hyalinisation in many cases.

John Milne. Aet 23 years. Ref. G.R.I. 3383:51. History. Swelling of neck present for past 16 months, not increasing in size. (His mother insists swelling was present before this).

The specimen is a large cervical lymph node 50 X 40 X 25 mm. of soft elastic consistency. It is difficult to decide the most appropriate category for this tumour. Under hand-lens magnification a distinct follicular pattern is visible, the false follicles being large, up to 3 mm. diameter. At higher magnification this pattern is lost and under the high power objective the tumour closely resembles cellular Hodgkin's disease. Reticulin compression is not a feature round the false follicles. Several small lymphoid follicles, some with pale centres can be distinguished at the periphery. The exoec growth phenomenon is demonstrated imperfectly. The capsule is irregularly thickened chiefly at the bases of trabeculae, it is intact and hyaline. Fibrosis is very tenuous but all forms can be seen. The reticulum cell increase is productive of immature lymphocytes and large morbid mononuclear cells. Hodgkin giant-cells can be identified in many places. Eosinophils are absent. Plasma cells are few and neutrophils are mainly concentrated in young granulation tissue round a large infarct. Mitoses average 1 per H.P. field.


History. Progressive asthenia for two years. Enlarged nodes not complained of but found on careful clinical examination. The present specimen is a node from the axilla. (Patient died at home six months later).

This is atypical Hodgkin's disease of fibroblastic type at a late stage. Initially this diagnosis was suggested by fibrosis only, many sections were prepared before the characteristic pathognomonic picture was found.

Normal lymphoid structure is totally lost. The capsular thickening is gross but the collagen is still in lamellae. Isolation nodules of the new tissue are enclosed in it, concentric re-formation is pronounced. The node is cut into a very coarse mosaic by enormous trabeculae. Fibrosis in all forms except coralline deposit is confirmed. In the cellular portion typical morbid mononuclear and Hodgkin giant-cells are present but most of the reticulum cell proliferation is expressed in the production of cells very close to typical fibroblasts. Eosinophils are still copious but their distribution in the node is characteristically capricious. Plasma cells are unusually numerous and fairly uniformly dispersed. Neutrophils are rare. Mitoses are very few indeed. Necrosis is not a feature.


History. Progressive asthenia for several months, there is generalised lymphadenopathy but the peripheral blood shows only secondary anaemia.

This is early lymphosarcoma in a cervical lymph node. There is an imperfect peripheral rim of surviving lymphoid tissue; in some localities pale centres are visible in it but the zones of lymphocytes surrounding them are particularly light. The exodic growth phenomenon is well shewn, for a neoplasm of this character. The capsule is of normal thickness but hyaline, infiltration is present at several points. Fibrosis is very slight, perivascular deposits and tenuous diffuse fibrils constitute most of it. The tumour cell is an immature lymphocyte, a free cell with a spheroidal nucleus which is larger and less chromatic than that of a normal small lymphocyte. Morbid mononuclear reticulum cells are very rare, Hodgkin giant-cells are absent. Eosinophils are absent but occasional neutrophils are seen. Plasma cells are scanty. Mitoses average 2-3 per H.P. field. Necrosis is not a feature.

Note. Tumour cells are visible in blood vessels and lymphatics.

Diagnosis. Lymphosarcoma, small cell type, early.

History.  Generalised lymphadenopathy with enlargement of the spleen.  The Wassermann reaction was negative.  The leucocyte count was 50,000 per cu.mm., the great majority were small lymphocytes.  The specimen is an axillary lymph node.

This is lymphosarcoma with leukaemia.  The tumour shews a moderately large morbid mononuclear reticulum cell component and fibrosis of the type seen in Hodgkin's disease is present.

The marginal sinus is preserved at some levels; an unusual finding.  The exodic growth phenomenon is apparent and the new tissue is disposed in a very faint nodular pattern which is best appreciated naked-eye.  The capsule is thickened and hyaline in remarkable degree but invasion with extracapsular deposit is present.  Trabeculae are present, they are thickened and partly hyaline.  Fibrosis of all types seen in Hodgkin's disease is confirmed even coralline deposits are present in quantity.  The type cells of the tumour are 1) an immature lymphocyte which preponderates, and 2) morbid mononuclear reticulum cells which are however rather small.  A very few typical Hodgkin giant-cells are present.  Eosinophils are absent and plasma cells are scanty.  Neutrophils are totally lacking.  Mitoses average 3 per H.P. field; necrosis is limited to individual scattered cells.

Diagnosis.  Lymphosarcoma, intermediate cell type.  Chronic lymphatic leukaemia present.  Affinities to Hodgkin's disease are remarkably close, especially noteworthy are fibrosis and reticulum cell content.

History. In the course of a surgical operation for the repair of an inguinal hernia on the right side this node, Cloquet's gland, was removed for good measure.

The node is relatively large but not unduly so considering the patient's age.

This is an example of extreme physiological hyperplasia which under different circumstances might justifiably prompt a diagnosis of tumour at an early stage. There is general retention of the normal architecture pattern but the pale centres of the peripheral germinal follicles are very large and some follicles are becoming diffuse. In some areas the edges of these merge imperceptibly with the pulp. The capsule is of normal thickness for this site. Trabeculae are poorly represented. Pathological fibrosis is not present - there is some fibrous tissue round the vessels from the hilar intrusion but this is typical of inguinal nodes. There is a marked increase of active reticulum cells both in the pale centres and in the pulp, large morbid mononuclear types are present but none have oxyphilic nucleoli. Giant-cells are absent. Eosinophils are very scanty, being well within normal. Plasma cells are increased above the normal number and often occur in small aggregates.

Curiously there is not any real sinus reaction.

Diagnosis. Physiological reaction, marked. Unusual follicular hyperplasia.
John McGuire. Aet 42 years. Ref. G.R.I. 298:52. History. Complaint of anorexia and loss of weight for the past 14 months. Clinical examination revealed generalised lymph node enlargement of moderate degree; the peripheral blood was normal. The patient received three courses of nitrogen mustard but died six months later (G.R.I. P.M.177:52). The initial biopsy, an axillary node, shews Brill's disease (follicular lymphoma). The necropsy material was poor but revealed total loss of follicular pattern and mutation to large cell lymphosarcoma.

In this specimen a thin mantle of normal small lymphocytes is seen at the periphery, the exodic growth phenomenon being discernible. The capsule is attenuated and split up by infiltrating small lymphocytes - the impression gained is that they are being driven through a disintegrating membrane. Concentric re-formation is visible in places. The new growth is disposed in false follicles of variable size, round these a little collagen is being irregularly deposited. The latter is the only form of fibrosis apparent. The proliferating reticulum cells of the pale centres are mainly shewing lymphoid differentiation but a few large morbid mononuclear types are seen. Hodgkin giant-cells are absent. Eosinophils, plasma cells, and neutrophils are conspicuously absent. In the necropsy specimen some uninvolved lymph nodes are present, e.g. deep cervical. In the remainder examined and in the spleen, the tumour can only be described as diffuse large cell lymphosarcoma. All trace of follicular pattern is obliterated and the type cell is a large primitive lymphoid series one with a leptochromatic vesicular nucleus.

/ Diagnosis.

History. Hard painless enlargement of the inguinal lymph nodes. The peripheral blood is normal.

This is Brill's disease in an inguinal lymph node. The tumour shows some invasion of the capsule.

There is not any survival of normal lymphoid tissue. The tumour is arranged in numerous large false follicles which are mutually contiguous and confluent in places. The capsule is imperfectly preserved, it appears thicker due to diffuse infiltration and expansion of the fibres by tumour cells. Trabeculae are poorly represented but where seen are unduly thick. Fibrosis of all forms is encountered but the amount is very small; perivascular lamellae account for most. (The collagen content is within normal for an inguinal node). The type cell of the growth is a small reticulum cell shewing lymphoid propensities. The nuclei are rarely spheroidal, most are indented and pachychromatic. Typical large morbid mononuclear reticulum cells and several small giant-cells are visible in certain fields. Eosinophils are absent. Plasma cells are occasional and neutrophils are restricted to one fragment of marginal sinus. Mitotic activity is high.

Diagnosis. Brill's disease (follicular lymphoma). Regarded as malignant, early loss of pattern.

History. A clinical diagnosis of primary gastric carcinoma was entertained. Surgical exploration revealed an externally normal stomach but enlarged hard lymph nodes in the lesser curvature. The lymph node recovered was originally diagnosed as late Hodgkin's disease but later linitis plastica (diffuse carcinoma) was established as the true diagnosis.

This node is widely replaced by maturing connective tissue and bears a remarkable resemblance to Hodgkin's disease. In this case meticulous examination of sections taken at four different levels revealed a few inconspicuous groups of contiguous epithelial cells and indicated the proper diagnosis. General features of help in this case included lack of hyaline fibrous thickening of the capsule, absence of trabeculae, a suspicious uniformity of the new connective tissue and persistence of sinus tissue. Eosinophils are lacking altogether, an unusual finding in gastric lymph nodes.

Diagnosis. Metastatic carcinoma, mimicking Hodgkin's disease.

(True diagnosis confirmed later).

History. Enlarged cervical lymph nodes of several months duration. In 1948 patient had a thyroidectomy, the gland was reported as hyperplastic. the blood state is not known.

This is indeterminate primary tumour, probably early lymphosarcoma, in a cervical lymph node. The diagnosis is seriously hindered by widespread avulsion of the capsule. There is an imperfect rind of normal hyperplastic lymphoid tissue at the periphery, in a few places small fragments of the marginal sinus can be identified. The exodic growth phenomenon is seen. Centrally the normal tissue is replaced by a very uniform lymphocytic growth. With very low power magnification a vague follicular pattern is discernible at one level. The capsule, where it exists is attenuated except round the inferent vessels. Concentric re-formation is absent. Trabeculae are very poorly represented. Fibrosis is minimal, there is merely hyalinisation of the vessels and these are scanty. The reticulin mesh however is slightly denser than normal. The type cell of the growth is a well formed slightly lepto-chromatic immature lymphocyte. Morbid mononuclear and rare mirror-image binucleate reticulum cells are evenly scattered through the tumour. All granulocytes and plasma cells are virtually absent. Mitoses average 2 per H.P. field. Necrosis is absent.

Diagnosis. Probably early small cell lymphosarcoma. Exodic growth phenomenon present.

This is a cervical lymph node 2 X 1 X 1 cm. which shows very early atypical Hodgkin's disease, the new tissue is almost all confined to the pulp.

There is considerable retention of normal structure; lymphoid follicles, some with pale centres, sinus tissue and parts of the pulp are preserved. The new tissue is patchy and chiefly in the medulla of the node, it reaches the capsule in one level examined. The lesion is too early to show the exodic growth phenomenon but incipient crushing of the cortex is seen at one point. The capsule is definitely thickened and this change affects the few trabeculae present as well. Infiltration is absent. Fibrosis is slight but definitely adequate to cause suspicion. Perivascular deposit, fibrils, and fibrillae are the forms present. Reticulum cell proliferation is manifest in the morbid pulp. The cells produced include perfectly typical morbid mononuclear types, Dorothy Reed mirror-image cells, and rare Hodgkin giant-cells. In addition small aggregates of epithelioid cells are common, some have large hyperchromic nuclei. Eosinophils are present but scanty, many are of lymphocytoid type. Plasma cells and neutrophils are readily found. Mitoses average 1-2 per H.P. field in the new tissue areas. Small focal necroses are common.

/ Diagnosis.

Note. The patient developed large hard lymph nodes all over the body and the anaemia never remitted. She became progressively asthenic and cachectic and died 26th October 1952. Necropsy was not allowed.
John Leather. Aet 37 years. A fishmonger. Ref. G.R.I. 1120:52, 1333:52. History. Generalised lymph node enlargement which has been in progress for three months. Abdominal pain has been present for one year, it is epigastric. The initial biopsy was badly mutilated and a repeat was requested, unfortunately an inguinal node was sent.

Both specimens are poor material, they were sent from Helensburgh and inevitably suffered autolysis due to delay.

The second specimen was reported carefully with an account of the changes seen and the diagnosis was left open. The alteration from normal is very slight. The changes giving rise to doubt are -

1. A very slight loss of general architecture with replacement especially in the deeper part of the node by indistinctly follicular lymphoid tissue.

2. A process of re-population of the atrophy fat-cone by lymphoid tissue, indicating proliferative activity.

3. Many individual cells both of the lymphocyte and reticulum series have unduly large nuclei.

The capsule and trabeculae are fibrous but the site (groin) precludes assessing this accurately. Fibrosis is appreciable round vessels and as fibrils and fibrillae in the pulp. Reticulum cell proliferation is present and though most cells are within normal, a few morbid mononuclear types are seen. Eosinophils are few, neutrophils and plasma cells are very low, too low to regard the lesion as inflammatory. Mitoses average less than 1 per H.P. field.

/diagnosis.
Diagnosis. Indeterminate lymphoid tissue hyperplasia, regarded as primary.

Note. This patient died of reticulum cell sarcoma within a year, which diagnosis was confirmed at necropsy (P.M.160:53). A large survey of lymph nodes from all groups shewed reticulum cell sarcoma of intermediate type. Deposits were also confirmed in the liver, spleen and bone marrow. A striking feature of this tumour was the degree of partial implication of the individual nodes.

History.  Swelling in the midline of the neck of indeterminate duration.

This is an example of probable physiological reactive change associated with marked fibrosis of the node capsule, trabeculae and part of the pulp.

There is general retention of normal active lymph node structure, follicles with pale centres are frequent.  The capsule is markedly thickened and hyaline, this change is uniform.  New perinodal lymphoid tissue, plasma cell deposits and etc. are absent from the specimen.  The trabeculae are abnormally thick in several instances.  Perivascular fibrosis is occasional and fibrils and fibrillae are visible in the pulp in one locality.  These latter appear the result of symplasmic transformation.  In several follicles the central arterioles shew fibrinoid degeneration, and perivascular macrophages with haemosiderin are seen.  The reticulum cells are active in the pale centres and in areas of the pulp, a few large, almost morbid varieties are present.  One giant-cell is seen at one level, it conforms to the Hodgkin type but is small.  Eosinophils and plasma cells are present but within the range of normality.  A few scattered neutrophils are present.  Sinus reaction of open type is notable in the marginal sinus.

Diagnosis.  Physiological reaction.  Fibrosis of capsule marked.
Slight fibrosis of pulp.  Fibrinoid degeneration present in some vessels.

History. Admitted as a surgical emergency, provisional diagnosis given as 'perforated peptic ulcer'. Considered to be suffering from acute leukaemia. A leucocyte count on admission yielded about 20,000 per cu.mm. The haematologists reported the finding of large mononuclear cells in the peripheral blood, these they considered lymphoblastic. Severe anaemia was present.

The specimen is an axillary lymph node. In general architecture a close mimicry of normal is seen; a peripheral wide zone of normal anatomical cortex exists but underlying this is a homogeneous looking mass of closely packed cells with large deep staining nuclei. The capsule is thick and hyaline, concentric re-formation is apparent, it is enhanced by the striking exodic growth phenomenon present. Well formed trabeculae are seen. Fibrosis of all types is present. Most is vascular (hyalinisation) or peri-vascular. The reticulum cells shew increase with a fair proportion of forms shewing lymphoid tendencies. Typical morbid mononuclear varieties and rare Hodgkin giant-cells are demonstrable. The eosinophils are increased above normal, they appear mature types. Neutrophils are fairly numerous throughout and plasma cells are increased.

Note. This patient died four weeks later; before death the haematological diagnosis was revised to acute myeloid leukaemia.

Diagnosis. The original diagnosis, Acute Hodgkin's disease was considered the nearest category, chiefly on account of fibrosis and cellular pleomorphism. May be regarded as lymph node in acute leukaemia.

History. A painless lump appeared in the left side of the neck eight months ago. It has gradually increased in size. It was resected after the provisional diagnosis of lipoma and presented as a fleshy lobulated encapsulated mass 2\(\frac{3}{4}\) X 1\(\frac{1}{2}\) X 1 inches.

This is atypical Hodgkin's disease. The aberrant features include lack of eosinophils and predominance of small cells; it has some affinity to lymphosarcoma.

Normal lymphoid tissue does not survive. The capsule shows very irregular thickening in places but the unaffected stretches are normal. There is doubtful infiltration at several points and concentric re-formation is visible. New trabeculae composed of swathes of immature connective tissue are forming, giving a coarse folliculoid pattern. Fibrosis is present in all forms. The reticulum cell proliferation tends to be focal, it is seen chiefly near the collagen deposits and near the edge. Large morbid mononuclear types and Hodgkin giant-cells are present but less florid than usual. Eosinophils are very sparse, small lymphocytoid cells with coarse oxyphilic granules are among them. Plasma cells and neutrophils are also scanty. Mitoses are very few, less than 1 per H.P. field.


History. Painless enlarged lymph node of neck, duration uncertain.

This is atypical Hodgkin's disease; fibrosis is less than usual, and eosinophils are practically absent.

Normal lymphoid tissue is absent. The capsule shews irregular thickening. Well marked re-formation is present in some sections, invasion is not seen. The trabeculae are thickened and small secondary branches segment the new tissue locally. Fibrosis exists in all forms, perivascular lamellation is the most conspicuous, other forms, especially fibrillary, are scanty. The reticulum cell proliferation is marked but tends to be focal. Large morbid mononuclear cells are common especially at the periphery of the new tissue. Hodgkin giant-cells are few, they are typical but small. Eosinophils are virtually absent. Plasma cells are very few, small lymphocytes are very numerous and appear mature. A few scattered neutrophils are visible. Mitoses average 2 per H.P. field.


History. This node was recovered from the lesser omentum at the pyloric end of the stomach. Gastrectomy had been performed for the relief of a benign chronic peptic ulcer.

The lesion is undoubtedly reactive change but it offers an interesting physiological mimicry of Hodgkin's disease. This is partly by virtue of the topical eosinophilia, a very common finding in para-enteronic nodes.

The general architecture is normal. The capsule shews irregular local thickening and is hyaline. Concentric re-formation is present in marked degree at some parts. Fibrosis adequate to suggest ordinary Hodgkin's disease is lacking, but perivascular deposit is unusual round individual vessels. There is general reticulum cell increase but true morbid varies are absent, here size of the nuclei is the determining factor. Eosinophils are numerous and often focal. The other cells are also fairly common. Sinus tissue is inconspicuous considering this is a gastric node.

Diagnosis. Physiological reaction shewing Hodgkin-like features in mild degree.

History. Nodule present in the occipital region of the neck for the past four weeks, it is painless. Received as 'lymph node'.

This is a cervical lymph node presenting focal epithelioid cell aggregates which show a tendency to suppurate. In addition there is pronounced capsular thickening and extreme physiological reaction in the pulp.

There is general retention of the normal architecture of the node but sinus tissue is obscure. The pale centres are irregular in outline and some are diffuse. Capsular fibrosis is marked and is being augmented by concentric lamellae from its inner aspect. The trabeculae are increased in size. Fibrosis is seen round a few vessels but is inconstant, there is slight local (sub-capsular) fibril and fibrillary collagen. The reticulum cells evidence a definite proliferation. This is chiefly expressed in the form of epithelioid cell islets which vaguely suggest tubercles. In some cases micro-abscesses are present centrally. In addition the pale cell and pulp reticulum cells show increase with fairly numerous morbid mononuclear forms. A few mirror image types are found but giant-cells are absent. Eosinophils are very few. Plasma cells are numerous and many are large.

Acid/alcohol fast bacilli cannot be found in sections appropriately stained.

Diagnosis. Physiological reactive change, possibly tuberculosis. Marked capsular fibrosis.

History. Swelling of the cervical lymph nodes of one year's duration. Recent swelling of the glands in both axillae. Old history of "chest infection". (There is not any active tuberculous disease radiologically).

This is typical Hodgkin's disease in a cervical node. A disrupted rind of normal lymphoid tissue is present below the capsule. The exodic growth phenomenon is present but disappearing. The new tissue is disposed in coarse irregular follicles. The capsule is grossly thickened and hyaline. Concentric re-formation is discernible; there is not any invasion. Fibrosis of all varieties is seen, the perivascular lamellation is remarkable. The reticulum cell proliferation is pronounced, large morbid mononuclear and typical Hodgkin giant-cells are common. Many of the latter are necrotic. Eosinophilia is intense. Plasma cells are recognisable with difficulty, the neutrophils are likewise hard to identify. Mitoses average 1 per H.P. field. Very small necrotic foci exist.

Diagnosis. Hodgkin's disease, typical.

History. Dysphagia had been present for six months. A sub-glottal extrinsic swelling was found on examination and there were firm matted lymph nodes present in both posterior triangles of the neck. Oesophagoscopv was negative.

This is an example of late atypical Hodgkin's disease, the growth is tumour-like. The specimen can no longer be identified as lymphadenoid. The cellular deposits of the new tissue are predominantly immature small lymphocyte masses and are set in a coarse hyaline collagenous mesh. Fibrosis of all forms is seen. Fibrils and fibrillae are relatively inconspicuous; the other forms are marked. The reticulum cell proliferation is still active but many of the cells produced are necrotic or going necrotic. Typical morbid mononuclear cells and Hodgkin giant-cells are readily found. A few atypical giant-cells like large foreign body giant-cells are visible, they appear to be fusion products of epithelioid cells. Several tiny foci of eosinophils exist but large areas are totally devoid of them. Plasma cells are numerous, especially at the edges of the cellular tissue. Neutrophils are virtually absent. Mitoses are very few.


History. Enlarged lymph nodes in both axillae, duration uncertain.

This is an axillary lymph node shewing lymphosarcoma arising from follicular lymphoma. There is a moderate tendency to sclerose, other minor affinities to Hodgkin's disease are apparent.

The entire node is morbid. The tumour shews a definite follicular pattern which is fairly uniform, the peripheral follicles are larger, becoming diffuse and shew the characteristic artefact shrinkage space round them. The capsule is widely infiltrated by tumour cells but persistent fragments are thick. Fibrosis of all forms can be found. Most is in the form of compacting fibrils round the false follicles, the other varieties are sparse and inconstant. The type cell of the growth is an immature (intermediate) lymphocyte but scanty morbid mononuclear reticulum cells and very rare small multinucleate giant-cells are also present. Eosinophils are present, mostly as scattered groups of 3-6. Plasma cells and neutrophils are rare. Mitoses are rare, about 1 per 2 H.P. fields. Necrosis is confined to occasional individual cells.

Thomas Quigley. Aet 59 years. Ref. G.R.I. 4910:52. History. (From Alexander Hospital, Coatbridge). A solitary glandular swelling appeared in the right axilla, it was discovered suddenly by the patient when it was already large. The chest is clear radiologically and the general health is good. Two nodes were sent, one 4 X 3 X 3 cm. the other 1 cm. diameter.

Both shew the same alterations. The lesion does not conform to any of the better characterised variants of lymphoid tissue tumour, it shews features of follicular lymphoma and Hodgkin's disease in about equal proportion. A rind of normal highly reactive lymphoid tissue is present in both nodes, it is 0.5 mm. wide. The exodic growth phenomenon is very distinct. The deep tissue is disposed in a definite but irregular follicular pattern. The capsule is not thickened, and it is intact. Light fibrosis is present. Perivascular lamellae, fibrils and fibrillae are identified. The reticulum cell proliferation is within the limits of normal but typical morbid mononuclear and true Hodgkin giant-cells are present. A few Langhan's type giant-cells are seen chiefly in groups of epithelioid cells. These latter aggregates are small and irregular, and unlike tuberculous reaction. Eosinophils are few. Plasma cells are very numerous, especially in the normal (reactive) peripheral tissue, and round the ill-defined deep follicles. Mitoses are rare. While the pattern is strongly suggestive of chronic irritation the nodes are suspiciously large.

Diagnosis. Lymphoid tissue tumour, relatively benign. Mingled features of Hodgkin's disease and follicular lymphoma present, but insufficiently characteristic to warrant diagnosis.

History. Asthenia of increasing severity over the last three years. A painful swelling appeared in the left side of the neck recently.

This is a cervical lymph node shewing a highly cellular pleiomorphic Hodgkin's disease picture; the appearances approach Hodgkin's sarcoma (type II of Warthin) but eosinophils persist, albeit reduced in number. Normal lymphoid tissue is absent. Concentric re-formation of the capsule is well shewn. The cellular part of the new tissue is irregularly folliculoid and set in a coarse collagen mesh. The capsule is grossly thickened and hyaline and permeated with islets of the new tissue. The appearances suggest that the perinodal compensatory lymphoid tissue has become implicated in the disease process. Fibrosis is very marked, all forms are represented but fibrillary deposits are few. The reticulum cell proliferation is florid and almost all are highly pleomorphic morbid types. Hodgkin giant-cells are very numerous. Rare Touton giant-cells are seen. Eosinophils persist but are scanty. Plasma cells are meagre, neutrophils are frequent including micro-abscesses. Mitoses average 2 per H.P. field in the cellular tissue. (It is likely that sepsis has supervened).


Note. This patient died in G.R.I. and necropsy was performed. See special P.M. series.

History. Asthenia and tiredness have been present for 15 months. During the past two weeks the left axillary nodes have been enlarged. The leucocyte count was 43,400 per cu.mm, the condition was regarded as subacute myeloid leukaemia.

This is a large axillary lymph node 4 X 3 X 3 cm. showing a remarkable picture of what may be termed acute or atavistic Hodgkin's disease; all the component cells are immature.

A few survival islets of small lymphocytes persist. Under low power the tumour appears uniform and cursory examination gives the impression of large cell lymphosarcoma. This is dispelled at once on scrutiny. The capsule is thick and transgressed at many points. Re-formation is not evident. Fibrosis is slight, tenuous deposits of all varieties are encountered. The commonest cell present is a spheroidal free cell with a large spheroidal or finely notched nucleus. Oxyphilic nucleoli are common and the nucleoplasm is deeper staining than that of typical morbid mononuclear cells. Often the cytoplasm is poorly defined and symplasmic habit obtains. Typical morbid mononuclear and rare Hodgkin giant-cells are identified. Eosinophils of myelocyte type and size are numerous, many lymphocytoid types are also seen. Plasma cells are very scanty. Many necrotic neutrophils are present especially at the sub-capsular periphery. Mitoses average 1-2 per H.P. field. Haemosiderin is present in the edge of the tumour.

Diagnosis. Atypical Hodgkin's disease, acute or atavistic type. Conspicuous de-differentiation of component cells.

/Note.
Note. This diagnosis was submitted before the blood state was established.

Revised diagnosis. Lymph node in myeloid leukaemia. The patient died two months later (P.M. G.R.I. 97:53). Generalised lymphadenopathy, splenomegaly and hepatomegaly were found; the femoral bone marrow shewed nodular white tissue of similar microscopical appearance.
James Loudon. Aet 25 years. Ref. G.R.I. 1460:53. History. An abscess developed in the left axilla approximately three months ago, the cause was obscure. It failed to resolve despite antibiotic therapy and a biopsy was taken on 3rd April. This was necrotic. This is a second specimen.

This node shews foci of reticulum cell sarcoma, and areas of typical Hodgkin's disease. The former lesion is a uniform stem-cell type of growth without the pleiomorphism of Hodgkin's sarcoma. Areas of normal lymph node tissue are present with active follicles well represented. Some deep sinuses and parts of the marginal sinus persist. The Hodgkin's disease areas are diffuse in the pulp; the reticulum cell sarcoma areas are better demarcated and some adjoin the former. The capsule is grossly thickened and hyaline, concentric re-formation is not seen but the exodic growth phenomenon is well shewn in places. Invasion is absent. Thick trabeculae are present. Fibrosis is present in all forms except coralline deposits, most is expressed as perivascular lamellation. The reticulum cell sarcoma is composed of large pale cells with indistinct (?) symplasma cytoplasmic outlines. The cytoplasm is feebly eosinophil. The nuclei are spheroidal with clear-line nuclear membranes and basophil granular nucleoplasm. Basophil and oxyphile nucleoli are common. Eosinophils are scattered through the deposits. In the Hodgkin's disease areas large morbid mononuclear types and Hodgkin giant-cells are present. Eosinophils are commoner in this tissue. Plasma cells and neutrophils are frequent in both growths.

Diagnosis. (Reticulum cell sarcoma, stem-cell type. (Hodgkin's disease, typical.

Co-exist in the same node but transition is difficult to trace.

Three lymph nodes from the groin, two large and one small. The largest measures 2.5 X 1 X 1.5 cm.

Two fairly distinct appearances are seen. The larger nodes reveal a soft cellular variety of Hodgkin's disease. These nodes do not contain any persisting normal tissue, the capsules are thin, even attenuated in places and are not much hyalinised. Invasion is not present and concentric re-formation is absent. Trabeculae are poorly represented. Fibrosis is slight, the chief expression is perivascular accretion, actual lamellation is rare. The vessels thus thickened produce a coarse imperfect segmentation of the new tissue. Diffuse fibrils and fibrillary deposits are present but inconspicuous because lymphoid series cells are still abundant. Occasional coralline deposits are noted. There is a characteristic increase in reticulum cells, most differentiating to the usual morbid type, but lymphoid differentiation is also seen. Many large mononuclear types and typical multinucleate giant-cells are present. Eosinophils are rare and plasma cells scanty. Neutrophils are also few. Mitoses are scarce, and necrosis is confined to occasional individual cells. The smallest node is normal in structure at one pole; the other part is expanded by a uniform cellular tumour of which the type-cell is the "owl-eye" variety of morbid mononuclear reticulum cell. Survival aggregates of lymphocytes are visible in interstices between the tumour cell masses. Eosinophils are absent and fibrosis is very slight indeed.

/Diagnosis.
Diagnosis.  
(1. Hodgkin's disease, atypical in so far as
   eosinophils are very scanty and fibrosis slight.
   Soft cellular and tumour-like.
(2. Reticulum cell sarcoma.

Note.  Patient died 1953.  Necropsy was performed, see special
P.W. series.

History. Vague illness of several months duration, generalised lymph node enlargement was found on examination. (Since diagnosis incipient chronic lymphatic leukaemia has developed).

This is a cervical lymph node shewing early small cell lymphosarcoma. Small scar-like areas of fibrosis are visible in the tumour. There is a beautifully preserved rind of hyperplastic lymphoid tissue approximately 0.5 mm wide between the tumour and capsule; it is not entire, the growth reaching the capsule at some levels. The exodic growth phenomenon is seldom better illustrated. The tumour is a very uniform mass of immature small lymphocytes. The capsule is attenuated, and is transgressed at several places. Trabeculae are absent. Fibrosis is limited to occasional perivascular deposits and small scar-like deposits. The latter are chiefly compacted fibrils. Reticulum cell nuclei including rare large morbid mononuclear forms with big nucleoli are easily found on scrutiny. Granulocytes and plasma cells are absent. Mitoses are numerous, up to 4 per H.P. field. Necrosis is not a feature.

Diagnosis. Small cell lymphosarcoma, incomplete involvement of node. Exodic growth phenomenon marked. Slight fibrosis.

History. Painless enlargement of lymph nodes in the left side of the neck and both axillae. Peripheral blood normal.

The specimen broke up during removal and the capsule became avulsed. Concentric re-formation is not seen. There is not any persistence of normal structure beyond a few survival islets of small lymphocytes. The trabeculae are very poorly represented. Fibrosis is meagre; perivascular lamellation and fibrillar deposits account for most, though coralline deposits are notable in one fragment. Reticulum cell proliferation is fairly uniform and more conspicuous than usual. Most cells of this line are morbid mononuclear types but macrophage types are also well represented. This pleiomorphism is pronounced. Hodgkin giant-cells are numerous, necrosis is common in individual examples. The eosinophils tend to be focal, those localities are in healthy Hodgkin tissue. Plasma cells are very few indeed, the lymphocytes are almost all rather larger than normal small lymphocytes. The growth rate appears about the usual.


There is not any survival of normal tissue but a few small aggregates of small lymphocytes are notable at the periphery. The exodic growth phenomenon is almost lost. The node capsule is hyalinised and much attenuated; at one point a definite breach is visible. There is much tumour outside the capsule at all points and it is infiltrating the fat. Fibrosis is early and slight, it is chiefly seen as fibrils and fibrillae, perivascular and coralline deposits are absent. (Senile hyaline degeneration is notable in the blood vessels). The node parenchyma consists of an active intermediate type reticulum cell sarcoma. Morbid mononuclear reticulum cells are the predominant cell but lymphocytic differentiation is also seen. Multinucleate giant-cells acceptable as Hodgkin giant-cells are scattered through the tumour. Eosinophils are absent, other granulocytes are uncommon, and plasma cells are scanty. Mitotic figures average 10-12 per high power field. Necrosis is beginning in the central portion of the tumour, many of the Karyorhetic nuclei resemble neutrophils but their true character is confirmed by use of the oil immersion lens.

Diagnosis. Reticulum cell sarcoma, intermediate type. Fields very close to Hodgkin's disease are discernible.
J.S. 3061:53. History. A swelling was first noticed in the right side
of the neck 12th July 1953. Painless. The node is from the anterior
triangle, others were present.

Survival follicles with pale centres, aggregates of normal lymphocytes
and large segments of the marginal lymph sinus persist. The exodic growth
phenomenon is shewn, the peripheral mantle consists of equivocally hyper-
plastic/neoplastic lymphoid tissue. The node capsule is irregularly
thickened but still shews many nuclei, invasion is not present. Swathes
of delicate young collagen segment the new tissue irregularly. Fibrosis
is early, all forms are present. Reticulum cell increase is marked,
many of the nuclei are still small and normal looking, but morbid mono-
nuclear varieties are present. Typical multinucleate giant-cells are
seen, they are more numerous in focal areas. Eosinophilia is pronounced
throughout, but plasma cells are scanty. Many of the lymphocytes present
are young forms with big nuclei. Neutrophils are occasional, micro-
abscesses are not seen. Mitoses average rather below 1 per H.P. field,
necrosis is not a feature.

Diagnosis. Hodgkin's disease, typical, early.

History. Has experienced pain in the back for several months and during the last month a swelling has appeared in the left side of the neck. Her general health is poor.

This is a cervical lymph node shewing Hodgkin's disease of the soft cellular type. Fibrosis is slight and the lesion is probably early. Several small poorly demarcated deposits of normal lymphoid tissue are seen at the periphery; some are surrounded by the new growth. Fragments of the marginal sinus persist. The new tissue is diffuse, nodulation is not appreciable. The capsule is thickened but hyalinisation is slight as yet. Eosinophils and plasma cells are frequent in between the compacting new lamellae. Trabeculae are not represented. Fibrosis is light; perivascular lamellae, fibrils, and fibrillae are the forms encountered. The reticulum cell overgrowth is marked, typical morbid mononuclear cells and Hodgkin giant-cells are frequent. Neutrophils are commoner than usual, and several micro- abscesses are present. Mitoses average 2 per H.P. field. Necrosis is not a feature.

Diagnosis. Hodgkin's disease, accept as typical; fairly soft cellular variety, sclerosis slight. Incomplete involvement of node.

Fragments of the marginal sinus and the deeper sinuses persist. Embedded deep in the smaller node are two minute islets of apparently normal lymphoid tissue, the lymphocytes here are distinctly smaller than in the neoplastic part. Very faintly marked follicular growth is apparent especially in the larger node. The capsules are hyalinised and attenuated, at several points they are over-run by tumour lymphocytes with the formation of extra-capsular deposits. Fibrosis is restricted being chiefly perivascular lamellation. Reticulum cells are scanty, most are peripheral and are already shewing lymphopoietic differentiation. Hodgkin giant-cells are not present, nor are there any eosinophils. The type cell of this tumour is an immature lymphocyte. Mitoses are relatively scanty but much variation is seen locally. There are two areas of necrosis in the smaller node and in the vicinity a few neutrophils are seen.

Note. Silver impregnation reveals an unusual amount of reticulin - enhancement of medullary mesh especially.

Diagnosis. Small celled lymphosarcoma with slight fibrosis, restricted to the vicinity of blood vessels.

Normal tissue persists in survival islets situated between the ill-defined follicles in which the tumour is growing. The capsule shows irregular thickening at the bases of trabeculae. The trabeculae proper are few, and form a scaffolding for new connective tissue swathes which imperfectly delineate the false follicles. Fibrosis is mainly of diffuse fibril and fibrillary character but occasional coralline deposits are seen. The proliferation of morbid reticulum cells is diffuse and unusually homogeneous, many typical mononuclear morbid types are present; Hodgkin giant-cells are present and are aggregated locally. Eosinophils are copious, particularly in the new connective tissue, plasma cells are very scanty. Neutrophils are unusually numerous and there is much focal necrosis. Mitoses do not exceed the usual number.


History. Five years previously patient developed a lump in the mid line of the sternum, biopsy was contemplated but not performed. She was treated outside when the lump became larger and ulcerated. This ulcer subsequently destroyed much of the skin of the breast and chest wall, and she complained of intermittent swellings in the neck. In July 1953 a biopsy of the edge of the ulcer was taken, ref. 2736:53.

(i) Skin. Due to gross sepsis with abscess formation, diagnosis was difficult. Follicular formations of tissue resembling Hodgkin's tissue were present; fibrosis was seen, but eosinophils were absent. Provisionally I reported this as "mycosis fungoides, probably Hodgkin's disease" but insisted that lymph node biopsy was essential.

(ii) Cervical lymph node. The tissue sent was scarred and ragged. Examination of one of the sections revealed it to be atypical Hodgkin's disease at a late stage but included in the mass were two physiologically active lymph nodes.

The capsules of the Hodgkin's tissue mass have lost their identity due to advanced fibrosis, the mass is probably a fusion product. Hyaline change and concentric re-formation are pronounced and the latter, together with thickened pre-existing trabeculae and swathes of new connective tissue cut the glands into segments. The intervening tissue is Hodgkin's tissue in which multinucleate giant-cells are unusually numerous. Fibrils, fibrillae, and coralline deposits of collagen are present in most areas. In one or two areas histiocytic reticulum cells
are seen and in a few the sclerosing reticulum cells are so uniform that they recall fibrosarcoma. Eosinophils are present and tend to occur in foci. Plasma cells and neutrophils are present, the latter chiefly related to necrotic foci which are common. Activity is higher than usual, the average mitoses per high power field being 2.

Diagnosis. Atypical Hodgkin's disease, almost Hodgkin's sarcoma.

The node is entirely neoplastic. The capsule shows a little local thickening but the lamellae are frequently disrupted by infiltration with immature lymphocytes. The exodic growth phenomenon is present. Trabeculae are poorly developed, they tend to be displaced obliquely. Perivascular fibrosis is appreciable but inconstant, other forms are slight. There are small false follicles in the deeper parts of the node and these show filamentous collagen between contiguous surfaces. The tumour is predominantly composed of morbid mononuclear reticulum cells which exhibit much mutual pleiomorphism. Typical Hodgkin giant-cells are present but scanty. Eosinophils are present but require close search. Plasma cells and neutrophils are conspicuously few. The growth rate, judged by mitoses is high, many fields contain 5 or more mitotic figures. Necrosis is limited to individual cells. Deposits of the growth are present in lymphatics. The adjacent fat is invaded.


All normal structure is lost. A vague follicular pattern is discernible, the chinks between the irregular nodules being filled with lymphatic and blood vessels containing lymphocytes and a few necrotic cells. The capsule where preserved shows gross hyaline thickening. Trabeculae are few but show fibrous thickening, especially at their bases. Fibrosis is well established. Perivascular deposits are notable and diffuse fibril formations of scar-like distribution are seen. The reticulum cell increase is not marked but characteristic large morbid mononuclear cells are readily found. Hodgkin giant-cells are few. Eosinophils are present but are not numerous. Plasma cells and neutrophils are scanty in most parts. The growth rate is about average - 1-2 mitoses per high power field. Necrosis is not present.

Diagnosis. Hodgkin's disease, typical. Early, fibrosing from the start, diffuse growth.
Dorothy Fox. Aet 27. Ref. G.R.I. 3669:53. History. Swellings have appeared in both sides of the neck and both axillae over the past four weeks. Clinically the size increase is still active.

This is a supraclavicular lymph node shewing the metamorphosis of Hodgkin's disease to Hodgkin's sarcoma. A crescentic deposit of normal hyperplastic lymphoid tissue is visible at one segment of the sub-capsular periphery. Follicles, some with pale centres, loose lymphatic tissue, and pieces of sinus survive. In the latter channels morbid reticulum cells are numerous. The exodic growth phenomenon is well shewn. The capsule is grossly thickened and hyaline, some concentric re-formation is discernible. Transgression is not present at the levels examined. Much fibrosis of all forms is seen. New trabeculae are forming from swathes of young connective tissue, and cut the node into segments. The reticulum cell proliferation is unusually florid with the production of copious irregularly pleiomorphic morbid types, Hodgkin giant-cells are very numerous. The latter often occur in aggregates. Eosinophils are still present chiefly as scattered small groups. Plasma cells are common, the neutrophils are copious near the focal necroses which are present. Mitoses average 2 per H.P. field.

Andrew Shearer. Aet 46 years. A tractor driver. Ref. G.R.I. W.3. 4057:53. History. He developed a painless swelling in the left groin one year ago. This has increased in size gradually and he now has hard masses palpable in the left iliac fossa, the right groin, in both sides of the neck, and axillae. General health has remained good.

The specimen is a large axillary node, 3 X 3 X 1.5 cm. A thin marginal rind of normal reactive lymphoid tissue is discernible with occasional pieces of sinus tissue. The exodic growth phenomenon is displayed. The capsule is of normal thickness in its greater part but at one locality is increased. In a few places concentric re-formation is detected. The new tissue occupies all the central part of the node, it displays a very faint follicular pattern, the individual false follicles being small, and dense centrally. Fibrosis of all forms is discernible but the amount is small. The commonest cell present is an intermediate reticulum cell type, but morbid mononuclear types are present and a few mirror image binucleate cells can be found. Hodgkin giant-cells are absent. Eosinophils and neutrophils are extremely hard to find but some plasma cells are visible. Mitoses average 2 per H.P. field.

Diagnosis. Brill's disease, becoming diffuse. Regarded as disseminated. Fields exist which are strongly reminiscent of Hodgkin's disease, especially near the fibrous deposits in the pulp.

History. Complaint of sudden pain in the right iliac fossa accompanied by sickness. Examination revealed local swelling. This is a deep pelvic node removed at operation.

This is a lymph node shewing active soft cellular Hodgkin's disease which is tumour-like in structure. The identity as lymph node is almost certain, lymphatics with valves are present in the capsule. Meagre crescents of reactive lymphoid tissue including follicles, some with pale centres are present near the inner aspect of the capsule. The exodic growth phenomenon is passing off but in several places the expanding nodules are driving towards the periphery. The new tissue is arranged in an indistinct folliculoid pattern. The capsule is much thickened but hyalinisation is restricted to the outer zone. Concentric re-formation is discernible, infiltration is not. Fibrosis is slight except for the capsular and trabecular components, the latter are composed of compacting fibrils. Fibrillar deposit is universal in the new tissue, the silver impregnation and van Gieson picture are closely similar. Perivascular lamellae and coralline deposit are less marked. The reticulum cell proliferation is extreme. The cells produced are highly pleiomorphic; epithelioid cells, morbid mononuclear types and Hodgkin giant-cells are all present in profusion. Eosinophils are distinctly scanty. Plasma cells are present in the usual numbers, neutrophils are copious round several infarct-like areas of necrosis which are present. Mitoses average 2-3 per H.P. field.

/ Diagnosis. 

Note. In March 1954 this patient developed nodules in the skin which showed reticulum cell sarcoma, eosinophils not present.

4420:53. History. Patient presented with a glandular swelling in the posterior triangle of the left side of the neck. It was painless and had been present six months. Examination revealed enlargement of the mediastinal nodes on the left side and a moderate neutrophile leucocytosis. Elsewhere the nodes were normal clinically and the spleen was not palpable. General condition was good.

This is atypical Hodgkin's disease which is apparently a slowly evolving type. The specimen consists of a small physiologically hyperplastic node and a lymphoid mass. The latter is organised as nodules of Hodgkin's tissue set in a coarse fibrous mesh. The nodules are irregular in size 1-2 mm. diameter and the mesh up to 1 mm. thick. Capsule is not represented but gross trabecular thickening is seen. Fibrosis is pronounced, it is seen in all forms, especially as compacted diffuse fibrils. Relatively little is seen in the cellular parts of the growth. The reticulum cell proliferation is only moderate and tends to be focal. Morbid mononuclear types and Hodgkin giant-cells are easily identified. Eosinophilia is slight but plasma cells are numerous. The latter occur often in dense foci in the peripheral part of the specimen. Neutrophils are sparse, most are seen in the few small necrotic foci which are present. Mitoses are not seen in many H.P. fields.


History. Enlarged painless lymph nodes have been present in the neck and both axillae for three months.

This is a large axillary lymph node shewing atypical Hodgkin's disease of a peculiar type. Persistence of normal structure is difficult to assess. Much apparently normal lymphoid tissue is present throughout the node and in one part, fragments of an atrophy fat-cone are preserved. This lymphoid tissue contains many small solid follicles, the arrangement being better indicated in the silver impregnation. The exodic growth phenomenon is absent. The new growth is patchy. Most of the reticulum cell overgrowth is seen in the locus of sinuses, but some is present in patches of the new tissue related to fibrosis deep in the node. Scattered morbid cells can be found in the pulp away from sinus tissue; typical morbid mononuclear and Hodgkin giant-cells are numerous. Transition from littoral cells can be traced. The capsule is much thickened but the inner layers are not yet hyalinised. Concentric re-formation is not present, and infiltration is not appreciable. Fibrosis is slight, most is expressed in perivascular lamellae and swathes of compacting fibrils in new trabeculae. Eosinophils are sparse. Plasma cells and neutrophils are fairly numerous, the latter being common round small necrotic foci which are present. Mitoses average 1 per H.P. field in the lesions.

Diagnosis. Atypical Hodgkin's disease. An unusual type. Eosinophils sparse. Morbid reticulum cells tending to be localised to sinus loci. Possible sinus origin in this case.

History. Patient has noticed swelling of the lymph nodes of the left side of the neck for six months. The enlargement is painless and there are no other symptoms. The nodes were firm and matted, rendering biopsy difficult.

The specimen is a cervical lymph node 18 X 14 X 10 mm. There is a very tenuous rind of hyperplastic lymphoid tissue at one point, it is crescentic and isolated between the true and false capsule, the latter is the result of concentric re-formation. The true capsule is irregularly thickened and becoming hyalinised, definite breaches have been made in it at several places with spread of the tumour into the external tissue. Fibrosis is conspicuous; all forms including direct symplasmic trans­formation are evident. The silver impregnation and van Gieson stained preparations coincide remarkably closely with the haematoxylin and eosin section. The reticulum cell proliferation is almost universal and productive of uniform cells. While the typical morbid mononuclear and Hodgkin giant-cells are easily found the majority are immature lymphocyte and intermediate types. Eosinophils are absent, plasma cells and neutro­phils are rare. Mitotic figures are rare, below 1 per H.P. field. Necrosis is limited to individual cells.

Diagnosis. Atypical Hodgkin's disease, tumour-like, regard as malignant (transgression of capsule etc.) No eosinophilia. Affinities to reticulum cell sarcoma and primitive lymphosarcoma.

History. Multiple swellings of the neck, axillae, and groins which have been present for four years. The specimen sent consists of three cervical lymph nodes 15-5 mm. diameter.

This is a relatively pure small cell lymphosarcoma which may have arisen from a follicular lymphoma since a pseudo-follicular pattern is discernible in the deeper parts of the nodes. Persistence of normal structure is doubtful, several reaches of what may be normal (reactive) marginal sinus are present. The exodic growth phenomenon is absent. The node substance is replaced by a mass of immature small lymphocytes which in the deeper parts are arranged in a poorly organised micro-follicular pattern. The centres of these follicles are appreciated by being slightly less densely packed. The capsule is generally thin but local thickening is observed at several points. Fibrosis is absent, several vessels shew hyalinisation but distinct perivascular lamellae are absent. The vast majority of cells are immature small lymphocytes. A very few morbid mononuclear reticulum cells are present but typical Hodgkin giant-cells are not found. Eosinophils, plasma cells and neutrophils are absent. Mitoses average 1 per H.P. field, necrosis is absent.

Diagnosis. Small cell lymphosarcoma, pure; some affinities to follicular lymphoma.

History. Swellings in the neck were noted first five years ago, these increased in size very slowly. On admission gross dropsy, cough and dyspnoea, and generalised lymphadenopathy with splenomegaly were present.

This is of doubtful category, it may be regarded as highly cellular tumour-like Hodgkin's disease and with equal justification as reticulum cell sarcoma. (Hodgkin's sarcoma). The specimen is a cervical lymph node 10 X 9 X 8 mm. At several points at the edge crescentic pieces of hyperplastic normal lymphoid tissue survive; they are simple loose lymphatic tissue without follicles. The exodic growth phenomenon is well seen. The new tissue occupies the remainder of the node and is of very uniform diffuse habit - a large single nodule. The capsule shews local thickening but is not hyalinised at these points although it is elsewhere. Concentric re-formation and infiltration are not present. Fibrosis is present in the new tissue in all forms but it is sparse. The reticulum cell proliferation is florid, the great majority of the cells being free with morbid single-lobe nuclei of "owl-eye" type. A few epithelioid cells are present, and Hodgkin giant-cells are very numerous (4 per H.P. field). Eosinophils are present but scanty. The remaining cells are merely occasionally encountered. Mitoses average 5 per H.P. field, necrosis is limited to individual cells.

Diagnosis. (Highly cellular tumour-like Hodgkin's disease, (or reticulum cell sarcoma with Hodgkin's giant-cells. (Hodgkin's sarcoma).

Note. Patient died 18th November 1953, see Special Necropsy 418:53. G.R.I.

These are axillary lymph nodes containing nodular foci of reticulum cell sarcoma of stem-cell type with some multinucleate giant cells. These tumour nodules are extensively necrotic centrally. Surrounding them the node tissue shows the changes of typical Hodgkin's disease and it in turn is enveloped in normal reactive lymphoid tissue. Peripherally normal reactive lymphoid tissue with follicles and occasional remnants of sinus tissue. The Hodgkin's disease areas are diffuse in the pulp adjoining the nodules of reticulum cell sarcoma. The exodic growth phenomenon is shewn by both tumours. The true node capsule is much thickened by lamellae of young connective tissue, hyalinisation is confined to the outermost parts. Concentric re-formation is appreciable. Infiltration is not seen. Fibrosis is chiefly fibrillar and fibrillary, the other forms are seen but are little in amount. The reticulum cell sarcoma is not sclerosing much. In the Hodgkin lesion the reticulum cells conform to the usual morbid types including typical Hodgkin giant-cells. Eosinophils are numerous, many are large myelocytic or lymphocytoid types; they are much fewer in the reticulum cell sarcoma. Neutrophils are frequent; most occur in relation to the large necrotic centres of the tumour nodules. Mitoses average 1 per H.P. field in the Hodgkin's lesion and 5-6 per H.P. field in the reticulum cell sarcoma. In the latter the type cells closely approximate to morbid mononuclear types and are free cells. Small giant cells are common.

/Diagnosis.
Diagnosis.  (Hodgkin's disease, typical.
(Reticulum cell sarcoma, stem-cell type, chiefly.

(A form of Hodgkin's sarcoma, note the disproportionate rate of growth of the latter part; the Hodgkin lesion is early since the whole node is not involved).

History. Swelling of the cervical lymph nodes for nine months, recently the axillary nodes have enlarged. Subcutaneous nodes are palpable across the clavicles, progressive dyspnoea and cough are developing.

The specimen is a cervical lymph node shewing Hodgkin's disease which is acceptable as typical although large cells are more numerous than usual. Some highly reactive but probably physiological lymphoid tissue persists. This is seen in a small adjacent node included in the specimen and in a tongue-shaped process in the morbid one. The exodic growth phenomenon is just discernible in a few places. The tumour exhibits a very faint folliculoid habit of growth. The capsule is irregularly thickened, the process is best seen at the base of the trabeculae, hyaline change is in progress. Concentric re-formation is hardly appreciable, there is not any infiltration of the capsule. Fibrosis is seen in all forms. The fibrillary component is very well illustrated. The reticulum cell proliferation is florid. The production of large morbid mononuclear types is above average. Hodgkin giant-cells are numerous, many are small. Eosinophilia is marked, a small proportion of lymphocytoid variety are identified. Plasma cells and neutrophils are scanty. The majority of the lymphocytes are immature with large nuclei. Mitoses average 1 per H.P. field. Necrotic foci are small and scanty.

Diagnosis. Hodgkin's disease, accept as typical; large cells rather above average, verging towards cellular type.
David Milne. Aet 53 years. Ref. Law 915:53. History. Swelling was first noted in the neck in April this year. Primary tumour was suspected and two biopsies were performed, neither of these has become available for study. The patient developed a further swelling in the right axilla recently. The general health is good.

This is an axillary lymph node shewing typical Hodgkin's disease at a late stage; suppuration is added. There is not any survival of normal structure. The growth is folliculoid, the cellular parts of the lesion being set in a matrix of maturing collagenous connective tissue. The capsule where it can be identified is grossly thickened to 1 mm. or more. Hyalinisation is incomplete and concentric re-formation is poorly shewn. Infiltration is not seen but there is fusion with an adjacent node. Fibrosis in all forms is marked. The reticulum cell proliferation is of the usual order; morbid mononuclear types and typical Hodgkin giant-cells are conspicuous. Eosinophilia is moderate, the cells occur in foci in both the cellular and fibrous components. Neutrophils are very numerous in micro-abscesses which abound, in these areas necrosis is notable. Plasma cells are rare. The lymphocytes are immature in many parts. Mitoses are of the usual order.

Diagnosis. Hodgkin's disease, typical, late. Folliculoid habit well shewn, micro-abscesses common.

History. Complaint of glandular swellings in the groins, axillae and neck, present for two months. In addition ulceration has developed round the anus. The Wassermann reaction is negative, X-ray of the chest is negative and Frei's test is negative.

This is a cervical lymph node shewing a soft cellular tumour-like atypical Hodgkin's disease. (The skin lesion is tuberculous ulceration in which the microbes were demonstrated).

Normal lymphoid tissue is absent and the exodic growth phenomenon is not seen. A patchy folliculoid organisation of the new tissue is discernible. The capsule is slightly but uniformly thickened. Hyaline change is early. Infiltration is doubtful, two small intra-capsular deposits of lymphoid tissue which are seen are almost certainly physiological. Fibrosis is slight. All forms can be made out except coralline. Most is still fine argyrophile fibrillary. The reticulum cell proliferation is marked and productive of many morbid mononuclear cells, Hodgkin giant-cells are meagre. Eosinophils are absent in significant numbers, neutrophils and plasma cells are rare. Necrosis is not a feature and mitoses are of the usual order.


History. Glandular swellings have been present in the neck for the past ten weeks. They are painless and are not tender on palpation. No other accessible nodes are enlarged.

The specimen consists of two cervical lymph nodes shewing identical changes. The lesion is probably nearest to atypical Hodgkin's disease but features of follicular lymphoma and lymphosarcoma are also displayed. Normal architecture and structure are almost entirely lost. The exodic growth phenomenon is not discernible. The node substance is replaced by a more or less well defined follicular formation in the interstices of which is tissue conforming to that of cellular Hodgkin's disease. In a few of the follicles normal pale centre cells are present but most are almost entirely composed of large immature lymphocyte nuclei, rare morbid reticulum cells are seen. The capsule is thickened fairly uniformly, infiltration by lymphocytes is seen, and some lymphatics are filled with these cells. Concentric re-formation is not a feature. Fibrosis is slight. Trabeculae are not represented to any extent. Diffuse fibrils and fibrillae are present but scanty, occasional vessels show very characteristic perivascular lamellation. Coralline deposit is not seen. The reticulum cell proliferation is notable only in the inter-follicular matrix. Morbid mononuclear and typical Hodgkin giant-cells are numerous. Eosinophils nearly all of blood type are copious in these areas. Plasma cells and neutrophils are noted. Mitoses average less than 1 per H.P. field. Necrosis is common in the giant cells.

/Diagnosis.

History. Presented clinically as a case of carcinoma of the right breast. The mamma was removed surgically. Naked eye a scirrhus was not found, instead numerous large crimson lymph nodes presented. Enquiry elicited that the patient was treated at home four months previously for glandular fever, diagnosed solely on clinical grounds. (At examination an enlarged node was found in the left axilla). The patient died after operation and necropsy was not obtained.

The absence of cancer was confirmed by microscopy. The clinical tumour is caused by a lymphoid tissue tumour. The tumour does not conform to those ordinarily recognised but is morphologically nearest to cellular Hodgkin's disease. All nodes show the same changes. Myeloid leukaemia is also a possibility. There is total loss of normal structure. The new growth occupies all the substance of the node. The capsule is much thickened by hyaline fibrosis, 1.0-1.5 mm. thick. It is not infiltrated and recalls the appearance of Hodgkin's disease vividly. Fibrosis is otherwise slight, all forms apart from coralline deposit can be found but the perivascular form is not typical. The reticulum cell proliferation is pronounced. Epithelioid cells are common, and next in frequency, morbid mononuclear types. Small Hodgkin giant-cells are identified. Eosinophils, including many lymphocytoid types, are very numerous. Plasma cells are also remarkably common. Neutrophils are sparse. Mitoses average 4-6 per H.P. field. Nearly all lymphocytes present are immature forms. Necrosis is rare and confined to individual cells.

/Diagnosis.
Diagnosis. Lymphoid tissue sarcoma, indeterminate category but nearest to cellular Hodgkin's disease. Eosinophilia pronounced, fibrosis slight, less capsular. Affinities to reticulum cell sarcoma with some lymphoid propensities. (Similar to myeloid leukaemia picture).

Note. Periglandular infiltration is seen in the breast, and abnormal white cells are also seen in some blood vessels. The circulating blood was not studied unfortunately and this diagnosis cannot be confirmed. In any event such gross enlargement of lymph nodes is not typical of myeloid leukaemia.

History. A painless slowly increasing swelling in the right side of the neck of some months duration. The swelling is slightly fluctuant. Naked-eye the cut surface of the node is a delicate café au lait tint and a faint follicular pattern is discernible in it.

This is typical Hodgkin's disease at an early stage. Persisting normal structures include lymphoid follicles, several with normal pale centres, and survival islets of lymphocytes. These are chiefly disposed peripherally in crescentic areas. The exodic growth phenomenon is well shewn. The new tissue is developing almost exclusively in the pulp of the node and is diffuse. The capsule is hyaline and doubtfully thickened, concentric re-formation is not appreciable at one level. Infiltration is absent. Fibrosis is slight. Trabecular thickening is visible and rare perivascular deposit can be found. Fibrils and fibrillae are discernible on close scrutiny especially in the van Gieson stained section and the silver impregnation, symplasmic transformation and coralline deposits are not seen. Reticulum cell proliferation is well shewn. Epithelioid and morbid mononuclear forms are readily found, the latter are highly characteristic. Hodgkin giant-cells are few. Eosinophils are very numerous. Plasma cells and neutrophils are readily found. One typical focus of necrosis is present at one level. Mitoses are few, below 1 per H.P. field.


History. Enlarged lymph nodes present in the right inguinal region for 15 months. No other lymph nodes palpable. She had a phlebitis of the right leg in 1950, it lasted a month.

This is part of a large node shewing early typical Hodgkin's disease. The growth is cellular and fibrosis is still slight. Persistent normal structure is identified but is fragmentary, one germinal follicle with a pale centre is seen together with remnants of marginal sinus, at the extreme sub-capsular edge. The exodic growth phenomenon is not convincingly detected. The capsule is still of normal thickness and cellularity. The new tissue is essentially diffuse in habit although a delicate irregular segmentation is detected with very low power study. Fibrosis is light; new compaction trabeculae of diffuse fibrils are appearing, they have no connection with the capsule. Fibrillae are scanty, likewise perivascular accretions. Symplasmic transformation is in process in patches. Coralline deposits are absent. The reticulum cell proliferation is even and productive of epithelioid, morbid mononuclear and Hodgkin giant-cells. Eosinophils are numerous but other granulocytes and plasma cells are rare. The lymphocytes are slightly immature. Mitoses average 1 per H.P. field, necrosis is absent.

David Hatton. Aet 47 years. Ref. Law Hospital 828:52.

History. For the past 15 months asthenia, profuse sweating, and loss of weight have become increasingly severe. Severe secondary anaemia was found and enlarged glands were discovered in the right groin.

This is an inguinal lymph node shewing atypical Hodgkin's disease; eosinophils are absent, and necrosis is unusually marked. The growth is distinctly tumour-like in some parts. Further features of interest are the presence of fat tissue in the node substance and evidence of fat atrophy, presumably previous to primary growth in the gland. Normal lymphoid tissue does not survive, and consequently the exodic growth phenomenon is absent. The new tissue is chiefly irregularly diffuse and micro-folliculoid. The capsule is atrophied to extinction at one pole due to previous fat atrophy. At this point the Hodgkin tissue reaches the fat line directly. Elsewhere thickening is doubtful. Fibrosis is chiefly in fibril and fibrillar form diffusely throughout the new tissue, symplasmic transformation is also seen. Perivascular lamellae and coralline deposits are scanty. The reticulum cell proliferation is productive of many different cell types. Epithelioid and morbid mononuclear varieties with transition forms are commonest, Hodgkin giant-cells are also conspicuous. Eosinophils are virtually absent, and plasma cells and neutrophils are rare. Lymphocytes of mature type are still the predominant cells in most areas. Mitoses number about 1 per H.P. field, or less. Massive necrosis of infarct type is marked, in addition individual cell death is very common throughout otherwise viable growth.


History. Since June of this year the patient has experienced general malaise and presents a painless swelling in the left side of the neck.

These are cervical lymph nodes shewing an uncommon variant of atypical Hodgkin's disease. There is very extensive production of fibroblastic cells. Small wedges and crescentic areas of reactive lymphoid tissue survive in both nodes. The exodic growth phenomenon is not evident however. The new tissue is disposed in coarsely segmented fashion with much maturing fibrosis round the more cellular nodules. The capsule is still fairly cellular but is grossly thickened, concentric re-formation and infiltration are absent. Fibrosis is pronounced, all forms except symplasmic transformation are visible. The reticulum cell proliferation is very widely given over to cells of fibroblastic characters. Epithelioid cells are rare, morbid mononuclear and Hodgkin cells tend to be aggregated in ill defined foci. Eosinophils are numerous, other granulocytes and plasma cells are commoner than usual and appear related to necrosis. Mitoses are impossible to assess in this preparation but are probably few. Necrotic foci of characteristic type are seen.


History. Multiple glandular swellings were discovered in the neck during investigation of bouts of vomiting and jaundice which have developed since February of this year. There is an old history of pulmonary tuberculosis.

This is a cervical lymph node shewing almost typical Hodgkin's disease, the features short of the ideal include very general and conspicuous early fibrosis, and very striking reduction of lymphocytes. Persistence of normal structure is seen. A small peripheral crescent of reactive lymphoid tissue with sinus tissue survives, in addition several lymphoid follicles with pale centres are seen at one point. The exodic growth phenomenon is present but not conspicuous. The tumour is distinctly, if faintly, follicular in habit. Capsular thickening is not impressive possibly due to artefact loss, concentric re-formation is absent. Fibrosis is widespread, diffuse and marked. New filamentous trabeculae are seen springing from the capsule. Perivascular deposit is slight. Diffuse fibrils and fibrillae constitute most of the new collagen, the other types are rare. The reticulum cell proliferation is chiefly productive of epithelioid cell types but morbid mononuclear and Hodgkin giant-cells also abound. Eosinophilia is of the usual order. Plasma cells and neutrophils are few. The reduction of small lymphocytes is striking. Mitoses average 1 per H.P. field, necrosis is restricted to occasional individual cells.


History. A swelling appeared in the left side of the neck in March 1953 (about two months ago). This was provisionally diagnosed as a salivary gland calculus and resected.

This is an almost pure reticulum cell sarcoma. The tumour is fibrosing and in some areas where this is more in evidence the picture of Hodgkin's disease is closely simulated. The specimen is almost certainly a lymph node, a marginal rind of small lymphocytes persists in places but otherwise all vestige of normal structure is absent. The tumour is very uniform and diffuse in most blocks; in one, scar-like areas of fibrosis and diminished cellularity mimic Hodgkin's disease. The capsule is slightly thickened and hyaline, concentric re-formation is absent and transgression is doubtful. Fibrosis is slight. Most is in the form of a delicate fibril and fibrillary filigree. Some swathes of compacting fibrils are seen, perivascular lamellae and coralline deposit are rare. The reticulum cell neoplasia is an almost pure small morbid mononuclear type, rare multinucleate types are encountered but their cytoplasm is deficient. Eosinophils are very rare, plasma cells and other granulocytes are virtually absent. Mitoses average 5 per H.P. field. Necrosis is fairly pronounced, the foci are poorly demarcated.

Diagnosis. Reticulum cell sarcoma, almost pure single cell type. Some affinities to Hodgkin's disease.

History. Swellings have been noticed in the neck for the past three months. Recently those on the left side have become larger and softer. A provisional diagnosis of tuberculous lymphadenitis was made. (After recognition of the true pathology a thorough search was made to discover the primary growth, this has so far eluded detection).

This is an example of metastatic carcinoma of squamous cell type in lymph nodes. In several blocks the similarity to Hodgkin's disease is remarkable. The isolated tumour cells mimic morbid reticulum cells to perfection and have evoked a copious diffuse fibroblastic reaction in their vicinity. In addition eosinophil infiltration is marked. There is an indistinct follicular pattern which also recalls the disease. In the remaining blocks, chiefly from the nodes on the left side shew easily recognisable deposits of squamous cell cancer.

APPENDIX B.

NECROPSY SERIES PROTOCOLS
NECROPSY SERIES PROTOCOLS

These are in the same order as in the text, Volume I.

p. 227. The sheets of the individual necropsy reports are sub-numbered.

Illustrations.

Volume III  Figs. 117-205

Volume IV  p. 61-87.

**Anamnesis.** The patient, a retired carter was admitted to the Eastern District Hospital, Glasgow, on 25th October 1949. He complained of swelling of the lower limbs and scrotum of about three months duration. Loss of weight, breathlessness, and anorexia were also present. During his stay in hospital numerous investigations were carried out but the results were negative or inconclusive. A tentative diagnosis of neoplasm of lung was considered because his condition was deteriorating rapidly and lymph nodes became palpable in the axillae. A slight irregular fever persisted throughout his time in hospital and on 15th November a leucocytosis of 33,200 per cu.mm. was recorded. Blood taken for culture on several occasions was sterile. Death occurred on 3rd December 1949.

**Report of Necropsy.**

**External.** Emaciation is present but is partly masked by dropsy.

**Reticulo-endothelial system.** Lymph nodes. There is generalized lymph-adenopathy. The cervical lymph nodes are numerous, slightly enlarged and very firm; at the base of the neck they are partly matted together. The cut surfaces are a prevailing greyish-white with yellowish-white bands running irregularly from the capsules so as to cut the surfaces into a mosaic. In both axillae the nodes are enlarged, hard, and discrete. When cut they reveal the same appearances as in the cervical group. The mediastinal nodes are grossly enlarged, the right inferior tracheo-bronchial gland measures 5 X 3 X 2 cm. They are hard and nodular. The cut surfaces are coarsely mottled grey and white. In the abdomen the nodes along the upper border of the pancreas and alongside the aorta are the most conspicuously enlarged. On section they are seen to have thin tenuous capsules and fleshy homogeneous yellowish-pink gland substance. The nodes of the mesenteric group are unduly numerous but not much larger than normal. In both groins the nodes are large, hard, and closely similar to the axillary group. The spleen is moderately enlarged and firm due to chronic venous congestion. The cut surface is pale crimson and homogeneous except for a few small rounded nodules of yellowish-white tissue deep in the pulp.

**Cardiovascular system.** A terminal sero-fibrinous pericarditis is present, the heart shews only mild brown atrophy. There is senile ectasia of the aorta.

**Respiratory system.** Sero-fibrinous effusion is notable in both pleural sacs, (left cavity 1200 cc, right cavity 500 cc.) Both lungs are slaty grey, small, and partly collapsed. In the right upper lobe several small ill-defined abscesses are noted. In the right eparterial bronchus the mucosa is nodular but not ulcerated.

**Alimentary system.** The oesophagus, stomach, and intestines are normal.
The liver is moderately enlarged and displays the nut-meg pattern of chronic venous congestion. There are several widely scattered deposits of firm white tissue about 1 cm. diameter throughout the viscus. They are more angular and less discrete than typical metastatic cancer. The remainder of this system is normal.

Genito-urinary system. A small series of aligned soft yellowish-white nodules 2 mm. in diameter are present in the mucosa of the bladder, near the urethral orifice. The kidneys, ureters and prostate gland appear normal.

Haemopoietic system. The marrow in the ribs is copious, pale red, and contains minute white nodules. In the cervical vertebrae similar appearances are seen. The red marrow of the femur extends down to within 12 cm. of the lower end, a few minute nodules are discernible.

Endocrine system. The thyroid gland is small and fibrous. The single parathyroid gland identified is normal, and the adrenal glands are normal.

Microscopical examination:
General description of the lesions encountered.

Late typical Hodgkin's disease with very extensive fibrosis and scanty eosinophils is restricted to certain lymph nodes and small fields of the prostate gland. In most sites the lesion corresponds to the 'Hodgkin's sarcoma' of Warthin. Transitions between the two pictures, which are regarded as simply different degrees of the same process, are evident. In the majority of the latter appearance the growth presents a poorly demarcated nodular habit. The tissue is made up of Hodgkin giant-cells in profusion, morbid mononuclear reticulum cells, fibroblasts, and fibrocytes. Interspersed between these aggregates is a variable number of small lymphocytes, neutrophils, and plasma cells. Eosinophils are absent. Collagen is generally conspicuous as a filigree of medium and tenuous fibrils. Several exceptions are noted; in the abdominal nodes fibrosis is scanty and the type cell is a morbid mononuclear variety, the appearances being relatively uniform. In a right cervical node a distinct whorling of fibroblastic cells is visible producing an appearance which recalls fibro-sarcoma. None of the sarcomatous lesions appear very rapidly growing. (There was not any antecedent biopsy in this case so that comparison is not possible). Microscopical evidence of tuberculous or other disease is lacking.

Distribution of the lesion. (See diagram).

Both processes are observed in nodes from the cervical, axillary, and inguinal regions. The mediastinal nodes also shew this mixture, and so does one large right pelvic node. In the abdomen all the retroperitoneal nodes examined and the mesenteric nodes shew the sarcomatous lesion. One node from the portal fissure is exceptional in that the
ordinary form of the disease co-exists. The bone marrow from one sample of rib shows the sarcomatous lesion, likewise a piece of cervical vertebral body. The femoral marrow is not involved. The picture of Hodgkin's sarcoma is seen in all the spleen and liver lesions examined. The adrenal glands are free of the tumour. Extra-nodal deposits of the sarcomatous form are confirmed in the bladder and prostate gland, the latter also exhibiting fields which would qualify for ordinary Hodgkin's disease. The lungs and kidneys examined by random blocks are free of either lesion.

**Diagnosis and Summary.**

(Hodgkin's disease.
(Hodgkin's sarcoma (Warthin type II).

The latter preponderates and appears to have arisen on a basis of the former. This case is interpreted as an instance of metamorphosis of Hodgkin's disease to Hodgkin's sarcoma. The latter is an highly pleomorphic tumour exhibiting a low degree of malignancy. Anasarca and hydrothorax are notable. Survival of normal lymphoid tissue is slight.
Agnes McNiven, aet 32 years. Ref. G.R.I. P.M.201:49.

Anamnesis. The patient was fairly well till 1945. At the end of this year she began to experience vague weakness. This hardly amounted to indisposition till early in 1948 when her strength failed and she was obliged to leave work. Her sister died in 1940 at the age of 18 of pulmonary tuberculosis, and she has two brothers at present suffering from the same disease. In view of this it was surmised that she might have the same complaint, but investigation revealed only enlarged lymph nodes in the mediastinum. She was admitted to Stobhill Hospital in May 1948. At this time a painful swelling appeared in the left side of the neck, and in the following month the right side also became affected. A cervical lymph node was removed for microscopical examination 12th October 1948 and was reported to shew the changes of very active Hodgkin's disease. (Ref. 0578:48. Serial 188). By December of this year both axillae presented swellings and cachexia set in. Deep X-ray therapy was administered with transient benefit; her clinical condition deteriorated steadily and she suffered considerable pain up till death on 5th July 1949.

Report of Necropsy. 6 hours post-mortem.

External. An extremely cachectic woman, the skin over the chest is brownish-grey and desquamating.

Reticulo-endothelial system. Lymph nodes. The superficial cervical lymph nodes are scanty small and hard, several in the deep groups near the spine are enlarged, firm, and white on section. Most of the axillary nodes are small and shotty but on both sides a few large indurated ones are present. Nodes could not be found in either antecubital fossa. The paratracheal nodes are enlarged, pale pink, and firm. In the mediastinum there are many large morbid lymph nodes, measuring up to 3 cm. in their long axes. Most show central caseation but are otherwise firm and white. The intra-pulmonary nodes at the lung hila are similarly diseased and spread of tumour-like tissue from them extends along the bronchi into the lungs. In the abdomen the mesenteric and retroperitoneal nodes are involved in the disease process. The nodes of the lesser omentum along the lesser curvature of the stomach are enlarged, firm, and pale pink. Small hard white nodules are visible along the mesenteric insertion of the small intestine giving a beaded appearance, to a lesser extent a similar condition is seen in the mesocolon. The nodes at the root of the mesentery appear normal and those along the upper border of the pancreas also seem natural. Below the level of the renal arteries the para-aortic lymph nodes are considerably enlarged and rather fleshy. The cut surfaces are pale pink. The groin lymph nodes are few but several large hard white specimens are present among those found. The popliteal nodes are very small, examples are present on both sides. Spleen. 200 g. Four discrete white tumour-like nodules sharply outlined by zones of congestion.
congestion are present. Their outlines are polygonal rather than round. The pulp is otherwise normal.

Bone marrow. Deposits of the new tissue are doubtfully present in the bodies of the lumbar vertebrae but none are seen in the right femoral marrow, ribs or sternum. The diploe of the skull is normal.

Cardiovascular system. The parietal pericardium is thickened especially at the base being up to 4 mm. thick. It is studded with white tumour nodules 1-2 mm. diameter. The sac contains 250 cc. of blood-stained fluid in which fibrin is copious. The heart, 180 g., shows brown atrophy. The aortic and main arteries are natural being almost devoid of atheroma lesions.

Respiratory system. Nodules of tumour-like white tissue, 1-1.5 mm. diameter, are scattered over the parietal layer of the pleura on both sides. They are particularly numerous over the diaphragm. Approximately 1500 cc. of serous fluid occupy the left cavity. Both lungs are small, partly collapsed, and congested. In the right main bronchus an ulcer 1 cm. diameter is encountered overlying a lymph node in the lung substance. A yellowish-grey slough occupies the lumen of this bronchus.

Alimentary system. The oesophagus is clear of actual invasion but is fixed by the enlarged para-tracheal nodes and the pleural nodules. Small tumour-like nodules are present in the peritoneum chiefly the diaphragmatic area; free fluid is slight. A small malignant ulcer is present in the stomach in the lesser curvature near the pylorus. The intestines are normal internally, peyer's patches and the solitary lymphoid follicles are not at all conspicuous. The liver, 1060 g., is small and terminally congested. Repetitive close slicing discloses several small (under 5 mm.) nodules of the new tissue. The gall bladder and ducts are natural, the pancreas is normal.

Genito-urinary system. Kidneys, left 50 g., right 80 g.; both glands are small but otherwise normal. The left ureter is healthy but the right tube is encircled by enlarged para-aortic (pelvic) lymph nodes. Dissection reveals actual growth in the ureter wall. The bladder, uterus, and adnexa are natural.

Endocrine system. The hypophysis cerebri is normal. The thyroid gland is small but of normal texture. Both adrenal glands are partly embedded in tumour tissue, dissection reveals small beads of white growth pressing into the cortices but no deep deposits. The thymus gland cannot be satisfactorily identified.

Central nervous system. The meninges are pale but otherwise normal. The brain (1160 g.) is small, pale, and wet, dissection does not reveal disease.

/Microscopical
Microscopical examination:

1. General description of the lesions encountered.

The lesion shows some variation from site to site. In the lymph nodes, the highly cellular pleomorphic picture with retention of eosinophils seen in the biopsy (Serial 188), is displayed. In the spleen and ureter the picture is closer to the typical disease. In the alimentary and pulmonary deposits tumour-like features are seen. Gradation between the extremes encountered is difficult to follow. This case is considerably complicated by widespread active fibro-caseous tuberculosis. The latter is present in the lungs, liver and spleen.

2. Distribution of the lesion.

The lymph nodes are involved with the exception of the popliteal specimens and one from the mesentery. Tuberculous disease interferes with identification of the process in some mediastinal and para-aortic nodes. Deposits of the new tissue are not identified in the bone marrow of the femur and sternum, gelatinous degeneration is marked. The spleen shows the lesion well, many of the fields show considerable haemosiderin deposit. Tuberculosis co-exists but generally in distinct foci. The spleniculus present does not display the disease. In the four blocks of liver examined the disease is not visible, active and obsolete tuberculosis constitute the nodules seen naked-eye.

Adrenal glands. One was examined, the gland is partly encircled by the new tissue growing in lymph nodes. At one point a wedge of the growth is embedded in the gland but intrinsic deposits are not seen.

The disease is confirmed in the duode no-gastric junction (gastric side), ureter, and juxta-intestinal nodules (at insertion of mesentery). Highly cellular foci of the disease are seen in the lungs.

Deposits of the disease are confirmed in the pericardium, pleura, and peritoneum.

Diagnosis and summary.

Hodgkin's disease showing transition to the morphological variant, Hodgkin's sarcoma. The transition is imperfect and apparently static since alteration from the biopsy is slight. Fibro-caseous tuberculosis co-exists and several normal lymph nodes survive.

The lesion in this case of atypical Hodgkin's disease shows enhanced pleiomorphism and cellularity. Tumour-like propensities exist, but are restricted and show little alteration after four years. An unusual feature is the location of the disease process in the ureter and serous membranes.

Hydrothorax and serous effusion of pericardial sac are present. The survival of lymphoid tissue is very slight.

Anamnesis. In 1943 the patient, who was then aged 31 years, developed painless enlarged lymph nodes in the left side of the neck. The history available records that a diagnosis of Hodgkin's disease was made at that time but it is not known whether this was substantiated by microscopy. Biopsy was not performed in this institution till 28th February 1946, Ref. 289:46 Serial 117, with this specimen the diagnosis was confirmed. He received deep X-ray therapy in March 1946, September 1949 and again in February 1950. Subsequently he was treated with nitrogen mustard and blood transfusion.

He was admitted to G.R.I. in mid-May 1950, extremely ill. At this time he had severe anaemia and leucopenia, despite further transfusions his condition deteriorated progressively till death on 10th July 1950.

Report of necropsy. 2 hours post-mortem.

External. An adult man of slight build. Moderate cachexia is present but is masked by dropsy.

Reticulo-endothelial system. 1. Lymph nodes. The superficial and deep cervical nodes are numerous, distinctly enlarged, and firm. Those at the root of the neck and thoracic inlet are the largest, being up to 2 cm. long. All are pale greyish-pink and the cut surfaces are fibrous and wet. In both axillae nodes are scanty, small, and firm; despite close search nodes cannot be recovered from either antecubital fossa. The mediastinal lymph nodes are enlarged, firm, and pale. They are matted together, and are difficult to separate. The inferior tracheo-bronchial nodes are conspicuously enlarged; the right gland measuring 5 x 3 x 3 cm. The adjacent areolar tissue is somewhat gelatinous due to oedema. In the abdomen the retro-peritoneal nodes, especially the para-aortic and pelvic groups, are enlarged to the size of broad beans. Most are pale greyish-pink and firm. Those along the upper border of the pancreas and in the lesser omentum also show enlargement, but the small intestinal mesenteric nodes are normal in number and size. In the groins the nodes are small and scar-like. Two small nodes are present in the popliteal fossae, one on each side.

2. Spleen (500 g.) The gland is held to the parietes by light adhesions. It is uniformly enlarged and only moderately firm. In the cut surface small suet-like deposits of new tissue are visible, set in a uniform crimson matrix.

3. Bone-marrow. In the sternum the tissue is widely fatty but several small white nodules are discernible. Tumour deposit is not visible in the lumbar and lower thoracic vertebral bodies. The marrow in the right femur presents a marled red and white pattern throughout, the entire

/cavity
cavity being thus occupied. At least part of the paler constituent is fat.

**Respiratory system.** Recent fibrinous pleurisy is notable on both sides, approximately 500 ml. of serous fluid occupy the cavities. Lungs, left 270 g., right 340 g. Both organs are partly collapsed, relatively bloodless and mottled greyish-white. In the right lower lobe adjacent to the heart there is an indurated zone of pulmonary tissue, it is in the form of a roughly circular disc 3 cm. diameter and 1 cm. thick. It presents at the pleural surface. The bronchi and trachea are natural.

**Alimentary system.** Sero-fibrinous ascites (1200 ml.) obtains. Large soft yellow gelatinous clots are present in the fluid. In the stomach a tumour-like growth 3 X 2.5 cm. is present in the submucosa of the lesser curvature 4 cm. from the pylorus. It closely resembles malignant ulcer but the underlying muscle is freely moveable. Several satellite nodules of white firm tissue surround it. The small and large intestine are not remarkable. Liver (1990 g.) Uniformly enlarged, pale lilac and glaucous. In the cut surface widely scattered suet-like deposits of new tissue are visible. The largest measures 2 cm. diameter. A fine cirrhosis is present. Gall bladder and ducts natural. Pancreas normal.

**Cardiovascular system.** Pericardium natural. Heart (255 g.) moderately atrophic, there is patchy fibrosis in the posterior wall of the left ventricle. The aorta and main arteries are visibly atheromatous.

**Genito—urinary system.** Kidneys, left 180 g., right 170 g., both glands are pale but otherwise normal. The ureters, bladder and prostate gland appear healthy.

**Endocrine system.** The hypophysis cerebri is normal in size and pink. The thymus is atrophied almost to extinction. The thyroid gland is much atrophied the lobes being leaf-like, the parathyroid glands (four recovered) are natural. The adrenal glands and testes are of normal appearance.

**Central nervous system.** The meninges are normal. The brain (1420 g.) is not remarkable except for pallor.

**Microscopical examination:**

**General description of the lesions encountered.**

In most situations the lesion is recognisable as Hodgkin's disease. Fibrosis of hyaline character is conspicuous especially in the capsules and the anatomical medullae of lymph nodes. Coralline deposits and compacted fibres constitute most, but vascular hyalinisation is also present, possibly the result of radiotherapy. Diminution of lymphocytes is notable, when extreme it imparts an unwonted homogeneity to the cellular
cellular parts of the deposits which then approximate closely to reticulum cell sarcoma. Eosinophils are very scanty. Haemosiderin is present in small amount in most sites.

Compared with the original biopsy the growth is much more tumour-like but the growth rate does not appear high. (See biopsy, Serial 117).

Evidence of tuberculous or other disease is lacking.

Distribution of the lesion. (See diagram).

All the lymph nodes examined shew the lesion except one node from the left popliteal fossa and several from the mesentery of the small intestine. Deposits of the new tissue are present in the femoral bone marrow, which is also the seat of diffuse light fibrous replacement. In the spleen the lesion is nearest to typical, minute foci of necrosis abound, and interstitial fibrosis is extensive. The liver lesions are confirmed, necrosis is common in the bigger deposits and the resemblance to round cell sarcoma is close. The adrenal glands and hypophysis cerebri are free of deposits.

Extra-nodal involvement is confirmed in the stomach. Here the growth is virtually an infiltrating reticulum cell sarcoma. The naked-eye impression of failure to infiltrate is erroneous, the muscle is being actively permeated and destroyed.

The lung lesion is less coherent than apparent naked-eye. It consists of partly coalescent perivascular and peribronchial deposits. The appearances are typical.

Diagnosis and summary.

Hodgkin's disease with extreme but incomplete involvement of the lymph nodes. The microscopical picture is altering in various sites towards reticulum cell sarcoma.

The initial lesion was slightly atypical Hodgkin's disease. The presumed duration is seven years, and yet normal lymph nodes are still present. Treated by irradiation and cytotoxic drugs it now presents a close similarity to reticulum cell sarcoma morphologically. Anasarca, hydrothorax and ascites are pronounced. The impression gained from the examination indicates that a very marked reduction of lymphoid tissue has been produced by the disease and by its treatment.
Anamnesis. Towards the end of 1947 the patient noticed a swelling in the left side of the neck, this was painless and was followed shortly after by similar swellings in the left axilla. In November of that year she was admitted and biopsy was performed (Ref. G.R.I. 2146:47, Serial 72). The diagnosis of Hodgkin's disease was established and X-ray therapy was given with a rapid satisfactory response. She remained well till July 1949 when the swellings recurred and a further course of irradiation was given. In June 1950 she was manifestly much weaker and complained of swelling in the epigastrium. This was found to be due to enlargement of the liver. X-ray therapy and the exhibition of nitrogen mustard produced regression of this tumour and she remained fairly comfortable till the end of November 1950. She was then re-admitted extremely ill and failed to respond to any therapy. She died 2nd December 1950.

Note. There is a record of a previous operation on the left axilla in 1937. I have been unable to trace whether this was a biopsy. There is not any record in this department of material from this patient at that date.

Report of Necropsy. 5½ hours post-mortem.

External. An adult woman of slight build. Marked cachexia is present but is partly hidden by dropsy.

Reticulo-endothelial system. 1. Lymph nodes. The cervical nodes are diminished in number, at least in recognisable form, and those identified are small and firm. The cut surfaces are pink and white. In both axillae scanty nodes of the same appearance are encountered. Despite careful search nodes cannot be found in either antecubital fossa nor in the epitrochlear regions. The nodes at the root of the neck and the paratracheal chains on both sides are augmented beyond the usual number. Several of these nodes are fibrous and pale but the larger ones are only moderately firm and reddish-brown. In the mediastinum the nodes are appreciably enlarged, but probably not increased in number. The pulmonary hilar nodes are normal-looking.

Throughout the abdomen the retro-peritoneal and pelvic nodes are small, discrete and firm. Their cut surfaces are moist and pink. The mesenteric groups appear almost normal. In the left groin several large nodes, up to 2 cm. long are noted, they are firm and pink. The cut surfaces are homogeneous and cream coloured. The remaining nodes in this site and in the right groin are small, firm, and scar-like. Their cut surfaces are white. Nodes cannot be found in either popliteal fossa.

2. Spleen (230 g.). The gland is tense and firm. Section shows moderate uniform engorgement and a solitary white tumour nodule 1.5 cm. diameter near the hilum. The deposit is angular and surrounded by a deep /bright
Bone Marrow. The marrow in the sternum and spine is uniformly pale pink, resorption of the cancellous bone is not seen, and tumour nodules are not visible.

In the right femur only a small amount of red marrow remains; it is confined to the extremities, most of the space is occupied by fat.

Alimentary System. A small shallow ulcer 6 X 2 mm. is present on the dorsum of the tongue. The oesophagus is normal. Ascites, (4000 ml.), is present; the fluid is faintly opalescent. The stomach and intestines are normal. The liver (2,500 g.) is increased in size and weight by virtue of a large, roughly spheroidal tumour mass, 13 X 12 X 10 cm., which occupies much of the right lobe. The growth is creamy-white at the periphery, with occasional red puncta; this zone is about 1 cm. wide and gives way to a pale green very oedematous yet firm centre having the texture of watered silk. Small satellite nodules of suet-like appearance surround it. Further similar nodules are encountered in the left lobe. The gall bladder and ducts are natural. The pancreas is well-preserved; it is normal.

Respiratory System. The system is normal except for the presence of a solitary nodule of firm pale fawn tissue in the left basal pleura.

Cardiovascular System. Pericardium, normal; the fluid is slightly increased. The heart (190 g.) shows brown atrophy. Atheroma is very slight throughout the arterial system.

Genito-urinary System. Kidneys, (left 180 g., right 155 g.). The glands show only slight ischaemic scarring. The ureters and bladder are normal. The uterus shows evidence of a previous parous state but involution is normal. The left ovary is partly replaced by an hard fibrous tumour nodule 1 cm. diameter; the right gland is natural.

Endocrine System. The thyroid gland is small but the texture is normal. Both adrenal glands present normal appearances.

Central Nervous System. (The head was not opened due to lack of permission).

Microscopical Examination:
1. General description of the lesions encountered.

The lesions include mature, almost totally hyalinised, deposits of Hodgkin's disease, and cellular tumour corresponding closely to reticulum cell sarcoma. The two growths appear distinct; it is not possible to trace convincing transitions in single specimens but some deposits present intermediate pictures. The sclerosed Hodgkin's disease is confined to
lymph nodes and the collagenisation is intensified by vascular
hyalinisation. In some nodes the identity of the lesion would be equi-
vocal, the persisting nuclei are so sparse. In the sarcomatous lesion
the growth is cellular but a degree of dissociation is detected, this is
enhanced by the profound reduction of lymphocytes. These necropsy
lesions are far more tumour-like than the biopsy specimen.

Evidence of tuberculous disease is absent.

2. Distribution of the lesions.

Lymph nodes. Obsolete, almost totally collagenised Hodgkin's
disease affects an axillary node on the left side and part of one on the
right. A single para-aortic mass of collagen believed to be a node is
also present. In the cervical, para-tracheal, mediastinal and groin nodes
the lesion is more cellular, in some instances recalling reticulum cell
sarcoma. In the cervical group two curious pictures are seen. In one
node, left cervical lymph node I, the appearances simulate Hodgkin's
disease but fibrosis and disorganisation of node architecture are lacking.
In another of this group, node II, a uniform overgrowth of morbid
reticulum cells is seen. This is accentuated by the depletion of lympho-
cytes. The node architecture is disorganised by fine diffuse fibrosis.
In the groin nodes on the left side a uniform tumour composed of morbid
mononuclear and multinucleate cells is found, this lesion is sclerosing.

Normal lymph nodes (shewing atrophy) are identified in the left
axilla and mesentery. Several presumed nodes were found to be fat or
condensed areolar tissue.

The femoral bone marrow is almost aplastic, the tissue is chiefly
haemorrhagic fat. Tumour is absent. In the spleen the lesion is
identified as rather diffuse reticulum cell sarcoma which retains a back-
ground of Hodgkin's disease. Fibrosis is marked. In the liver the
growth is extensively necrotic but where the cellular tissue survives it
is composed of morbid reticulum cells with densely hyperchromic nuclei,
pleiomorphism is restricted.

The adrenal glands are natural.

Extra-nodal deposits of the tumour are identified in the left ovary
and in the pleural nodule noted. Rare tumour cells are found also in the
connective tissue of the pancreas. The lingual ulcer recorded is a
benign non-specific lesion.

Note. In one of the paratracheal nodes an unusual type of multi-
nucleate giant-cell is encountered, viz. a Touton giant-cell.

Diagnosis and summary.

Hodgkin's disease and reticulum cell sarcoma. While it is highly

/probable
probable the latter has supervened on the former it is not clearly
demonstrated. The virtual disappearance of lymphocytes is notable in the
hepatic and groin node lesions as well as in the late Hodgkin's disease
affected lymph nodes. This rather supports the view that these lesions
are related.

In this case the lymph node lesions as a whole are not so pronounced
as usual, this may be the result of therapy. Systematisation is
incomplete, but surviving lymphoid tissue is small in amount. Anasarca
and ascites are prominent features.
Andrew Young, aet 27 years. Ref. E.35:51.

Anamnesis. The patient was in good health till early in 1942 when, at the age of 18 he developed a painless swelling in the left side of the neck. This was regarded as tuberculous in character and he was treated for three months in Ruchill Hospital. Later in that year the swelling increased and he was admitted to Stobhill Hospital where biopsy established a diagnosis of Hodgkin's disease, Ref. Stobhill Hospital P42/834. Serial 57. Deep X-ray therapy was administered immediately afterwards and at intervals till 1945 when further nodes appeared in the left axilla. After a course of X-ray treatment given to destroy these, the patient developed numerous crops of boils which were very troublesome. The disease appeared to remit when the supplicative episodes ceased and he remained tolerably well till 1949 when symptoms recurred. Nitrogen mustard was administered and considerable improvement followed till the following year when the neck glands were grossly swollen. X-ray treatment was given but the response was poor. He became cachectic and bedridden from the beginning of September 1950, and died 21st February 1951.

Report of Necropsy. 6 hours post-mortem.

External. A tall well-developed man. Wasting and dropsy are evident in the limbs but the abdomen is full. The skin is a very pale cafe-au-lait tint, several warts are noted on the trunk.

Reticulo-endothelial system. 1. Lymph nodes. The cervical lymph nodes particularly those on the left side are very much increased in number and many are enlarged. They remain discrete in most cases, but the investing areolar tissue is much indurated. These nodes are extremely hard being almost cartilaginous in consistency. Their cut surfaces are white and faintly nodular. Small yellow areas of necrosis are seen in a few of the larger nodes. In the axillae the nodes are small scanty and hard. They are distinctly yellowish-grey. Nodes are not recoverable from either antecubital fossa. The mediastinal nodes are conspicuously numerous and large. They form a coarsely nodular coherent mass 12 X 9 X 8 cm. which encircles the lower half of the trachea. The individual nodes vary much in size. The smallest, the inferior group members measure 2-3 mm. in diameter whereas the largest, those above the lung roots measure 2-5 cm. diameter. They are all extremely hard, white, and nodular. Rarely small necrotic foci are encountered in them. Stibbe's nodes are much enlarged too, they are increased in number and are fused with a tumour-like plaque attached to the basal pericardium. Nodes in both lung hila are also involved by the disease. In the abdomen both the mesenteric and retro-peritoneal nodes are extensively diseased. In the lesser and greater omenta many large hard nodes are encountered. In the mesentery of the small intestine a mass of semi-fused nodes, 20 X 15 X 7 cm., occupies the root of that structure, and minute firm nodules often in small clusters are copious along the insertion of it into the bowel.
Several of the root nodes are enormous, up to 10 X 6 X 4 cm., occasional massive necrosis is a feature in these giant specimens. The para-aortic group form a roughly fusiform mass 11 X 7 X 4 cm., the individual nodes closely resemble those in the mesentery. In both groins scanty small hard white nodes are found, none could be recovered from the popliteal fossae.

2. Spleen. (450 g.). Uniformly enlarged, smooth and dull plum colour. Repeated close slicing and scrutiny fail to reveal any deposits of Hodgkin tissue. The appearances are simply those of engorgement.

3. Bone marrow. In the sternum ribs and spine the marrow is copious homogeneous and rich red, nodules are not visible. In the femur the marrow is pale yellow and gelatinous throughout, focal lesions are not present.

Cardiovascular system. Pericardium. There is tumour-like invasion of the parietal wall at the base of the heart, nodules of very hard white tissue 2-3 mm. diameter are visible from the internal surface. The heart (200 g.) shows brown atrophy. Atheroma is notably mild in the arterial system.

Respiratory system. Pleurae. Approximately 2000 ml. of pale straw-coloured fluid are present in the cavities. On the left side a dense fibrous pleurisy up to 3 mm. thick affects both layers of the membrane. In places fusion, with encystment of gelatinous pale yellow exudate, is noted. On the right side similar changes obtain but are less developed and are restricted to the upper lobe area. At the lower end of the trachea intra-mucosal or submucosal beaded growth of white tumour extends over an area 7 X 5 cm. The deposits show occasional colasence to form raised plaques, this growth extends down into the bronchi. Both lungs are partly collapsed. Nodules of hard white tissue are present near the hila of both. Some suppuration is detected in the left organ, apparently due to retention pneumonia. The diaphragm is patchily encrusted with deposits of hard white tissue.

Alimentary system. The oesophagus is normal. The stomach and intestines are very well preserved and internally they are normal. Peyer’s patches are very indistinct. The liver is normal apart from terminal congestion, repeated close slicing fails to reveal any tumour deposit. The gall bladder and ducts are natural. The pancreas is normal.

Genito-urinary system. The kidneys (left 160 g., right 150 g.) are normal. The ureters, bladder, prostate gland, and testes are natural.

Endocrine system. The hypophysis cerebri is normal. The thyroid gland is small but of normal texture, the two parathyroid glands recovered are natural. Both adrenal glands are normal apart from a curious dull olive and mahogany coloured mottling of the cortices.

Central nervous system. The meninges are healthy, the brain is normal in all respects.

/Microscopical
Microscopical examination:

1. General description of the lesions encountered.

Atypical Hodgkin’s disease, corresponding closely to the variety seen in the biopsy, only mature and extensively sclerosed, co-exists with a cellular tumour which is virtually Hodgkin’s sarcoma. Paradoxically transition between the two is imperceptible, they are so very similar. In the "sarcomatous" form the multinucleate giant-cells are very numerous indeed and sclerosis is less evident, the two pictures are simply different degrees of the same process. An unusual feature is the retention of eosinophils but these cells are nevertheless very scanty.

Suppurative broncho-pneumonia and organizing pleurisy are seen in the lungs but evidence of tuberculous disease is absent.

2. Distribution of the lesions. (See diagram).

All the lymph nodes examined show evidence of the disease in one or both its forms. The cervical, axillary and mediastinal nodes show chiefly the sarcomatous picture but fields of the densely fibrotic type are encountered in the last two places. In the abdomen the mesenteric and retro-peritoneal nodes reveal almost identical appearances with superior frequency of the sarcomatous lesion. In the groin nodes the less cellular sclerosing form predominates.

The samples of femoral bone marrow, liver, and spleen are free of the disease in either form. Both adrenal glands are unaffected.

Deposits of the new tissue are confirmed in the trachea, both lungs, the diaphragm, and the pleura. Both kidneys, the prostate gland and brain are unaffected. A sample of colon examined shows small normal lymphoid follicles in the submucosa.

Diagnosis and summary.

Atypical Hodgkin’s disease morphologically close to reticulum cell sarcoma, showing fairly complete metamorphosis to the morphological variant, Hodgkin’s sarcoma. The lesion, initially typified by unusual aggregation of the proliferating morbid reticulum cells shows enhancement of this phenomenon which together with remarkable production of multinucleate giant-cells gives the picture of Warthin’s reticulo-endothelio-blastoma. The lesion is not systematised, despite the elapse of nearly nine years. The pulmonary, tracheal, diaphragmatic, and pericardial deposits are probably autochthonous, though it must be conceded that the first named might be embolic in origin.

Oedema is notable in this case. Surviving normal lymphoid tissue is minimal but recognisable in the para-enteronic loci.

Anamnesis. The patient was well till 18 months ago when she began to feel weak and breathless. These symptoms intensified gradually, and dysphagia with vomiting became troublesome. Pulmonary tuberculosis was suspected and she was treated in Robroyston Hospital for some months, definite evidence of tuberculous infection was not however established. She was admitted to Glasgow Royal Infirmary 13th July 1950. At this time she was pale, sallow, and dropsical. A large pleural effusion, and enlargement of the spleen were discovered, but lymph node enlargement was not a feature. Considerable anaemia and moderate leucocytosis were present. Guinea pig inoculation of pleural fluid failed to incriminate Myco. tuberculosis as a cause. A diagnosis of "reticulosis, type uncertain" was considered and she received radiotherapy. She was then allowed home. She was re-admitted in October 1950 and spent nearly three months in hospital but her condition slowly deteriorated. On 31st January 1951 she was admitted again, manifestly in her last illness. She died 16th February 1951.

Note. It is established that lymph node biopsy was not performed in this case.

Report of Necropsy. 24 hours post-mortem.

External. A very emaciated young adult woman.

Reticulo-endothelial system. Lymph nodes. The cervical nodes are small and inconspicuous above the root of the neck. At the thoracic inlet several fleshy, moderately firm, glands are noted, these measure up to 2 cm. in their long axes. Their cut surfaces are pale pink and white. The axillary nodes are few in number on both sides but several large firm specimens exist. On section they are white and coarsely nodular. Nodes cannot be recovered from either antecubital fossa. The para-tracheal chains are prominent, the individual nodes varying from 1-3 cm. diameter. These glands are mutually separable with ease. In the mediastinum the lymph nodes are increased above the usual number and most are appreciably enlarged. On section they appear white except for scattered carbon particles. Stibbe's para-sternal nodes are readily seen due to their enlargement.

In the abdomen the retro-peritoneal nodes are numerous and enlarged. The most conspicuous are the para-aortic groups round the coeliac and renal arteries. The glands of the pelvis are very scanty and small. The mesenteric groups are normal in appearance. A solitary small lymph node is found in the left groin but none can be recovered from the right. Both popliteal fossae are similarly devoid of nodes.

Spleen. (190 g.). Three pale nodules are visible from the surface shining through the capsule. Dissection reveals these to be typical suet-
like masses of Hodgkin tissue approximately 1 cm. diameter.

The bone marrow of the right femur, lumbar and lower thoracic vertebrae, and ribs is pale red and homogeneous. Visible deposits in it are absent.

Respiratory system. 1200 cc. of greenish-yellow opalescent fluid lie in the right chest. The right lung is collapsed and covered by a thick fibrin deposit. Section reveals white tumour-like infiltration of the peribronchial tissue which extends out to involve most of the upper lobe. Necrosis is marked in the centres of the coarsely lobular deposits. Small islets of surviving lung tissue are enclosed by the growth and are also visible at the periphery. On both sides the bronchi are normal internally as far as dissection can be carried. The left lung is clear of disease, the only feature of note being a patch of fibrinous pleurisy.

Cardiovascular system. The pericardium is infiltrated in both its layers at the base, the new growth is hard, white, and nodular. Effusion to the extent of 500 cc. of faintly turbid fluid is noted. The heart 220 g. shews only atrophy. The aorta is natural, atheroma is minimal.

Genito-urinary system. The kidneys (left 145 g., right 145 g.) are normal. The ureters and bladder are healthy. The uterus and adnexa are natural.

Alimentary system. Ascites (1200 cc.) is present, fibrin is very scanty. The oesophagus, stomach, and intestines shew only post-mortem change. The liver (1720 g.) is large, a nut-meg pattern is discernible in the cut surface; despite repeated close slicing tumour deposit is not seen. The gall bladder and ducts are normal. The pancreas is natural.

Central nervous system. Brain (1320 g.) The organ is normal to naked-eye examination. The meninges are pale but otherwise normal.

Endocrine system. The hypophysis cerebri is small, red, and very soft. The thyroid gland is natural but rather small. An enlarged lymph node is partly embedded in the cortex of the right adrenal gland, the left gland is normal.

Microscopical examination:
1. General description of the lesions encountered.

The basic lesion is ageing Hodgkin's disease. This is seen in lymph nodes and the spleen. In these sites fibrosis is pronounced and infarct-like areas of necrosis are present in some cases. Eosinophils are retained in these deposits. In addition transition to the variant here termed Hodgkin's sarcoma is notable. In certain nodes examined both pictures can be seen, in others only the latter is exhibited. (Comparison with antecedent biopsy findings is not applicable).

/There
There is not any evidence microscopically of co-existent tuberculous or other disease. Preservation of the histological material is unfortunately not very good and this has limited successful photographic illustration.

2. Distribution of the lesions.

Lymph nodes. In the cervical, axillary, mediastinal, and three retroperitoneal nodes the picture is one of mature Hodgkin's disease. In one of Stibbe's nodes from the right side Hodgkin's sarcoma is seen in addition. The mesenteric nodes and several retro-peritoneal specimens shew only Hodgkin's sarcoma. Incomplete involvement of nodes is present in two specimens, one from the mesentery and in one para-aortic node. The node from the left groin is normal apart from moderate atrophy.

Only the femoral bone marrow was examined, neither lesion is identified in it. In the spleen the form is that of late cellular Hodgkin's disease, necrosis is slight but some haemosiderin deposit is visible. Growth of Hodgkin tissue of moderately fibrous type is confirmed round the right adrenal gland. The gland is closely invested but not penetrated, the new tissue is not present in the gland itself. Fat spaces in the new growth suggest that fatty tissue has been infiltrated in tumour-like fashion. The hypophysis cerebri is very poorly preserved, there is not any evidence of the new tissue in it.

Extra nodal sites shewing the disease include the pericardium and superficial myocardium. The cardiac muscle is being infiltrated by typical Hodgkin's disease. The right lung lesion is Hodgkin's disease. It assumes the form commonly seen namely a type of chronic interstitial pneumonia with a central necrotic zone. The disease is not found in any of the blocks of liver examined.

Diagnosis and summary.

Hodgkin's disease with metamorphosis to Hodgkin's sarcoma. The transition is displayed fairly convincingly in two sites. The tumour-like propensities of Hodgkin's disease are well shewn by the infiltrative tendencies seen in the heart and peri-adrenal fat. Failure of the disease to systematise is noteworthy. Hydrothorax, pericardial effusion, and ascites are pronounced. There is very severe atrophy of lymphoid tissue but some normal is identified.
David Hatton, aet 47 years. Ref. P.M. 131:52. Law Hospital.

Anamnesis. This patient was first admitted to Law Hospital on 12th August 1952. He complained of general weakness, profuse sweating (especially at night), loss of weight, and anorexia. These symptoms had been present for one year. He had remittent pyrexia and severe secondary anaemia. His condition improved after blood transfusion and he was transferred to Strathclyde Hospital. On 21st November 1952 he was re-admitted to Law Hospital where the Murchison phenomenon was observed during his sojourn. Enlarged lymph nodes were found in the right iliac fossa and groin. Biopsy was performed (Ref. Law Hospital 828:52., Serial 218). This showed Hodgkin's disease. His condition deteriorated rapidly, he became mentally confused and jaundice developed. He died 27th December 1952.

Report of Necropsy. 2 days post-mortem.

External. An extremely cachectic middle-aged man. Icterus is present. Healing surgical wounds are present in the right inguinal region, a para-median abdominal scar is also present.

Reticulo-endothelial system. 1. Lymph nodes. The cervical lymph nodes are normal in number and size but are appreciably firmer than usual. In the left axilla the nodes are enlarged, hard, and numerically increased, the largest measures 2.5 cm. diameter. On the right side they appear almost normal but are firm. The mediastinal and hilar nodes shew moderate anthracosis but are not enlarged nor increased in number. Their consistency is firm. In the abdomen the mesenteric and retro-peritoneal groups are involved in irregular fashion. Of the former those in the lesser omentum (porta hepatis), the transverse mesocolon, and coeliac group are enlarged. The biggest measure 3 cm. diameter. The nodes in the small intestine mesentery are small and inconspicuous. The para-aortic groups on both sides are considerably enlarged and firm. In the right side of the pelvis nodes are prominent along the right internal iliac vessels. The nodes in the left groin are small, fairly hard, and scanty, those in the right region are enlarged and firm. Nodes are present in both antecubital fossae and popliteal fossae, they are very small.

2. Spleen. 350 g. The gland is enlarged to over twice the normal size. It is firm and palpably nodular. The cut surface discloses some half-dozen discrete greyish-pink nodules of between 1-7 cm. diameter set in a dark crimson pulp. These tumours are not very hard.

3. Bone marrow. In the sternum the marrow is red and semi-fluid. The marrow of the right femur is greyish-green and gelatinous throughout.

Cardio-vascular system. Pericardium, natural. Heart (230 g.). Brown atrophy is present, the coronary arteries are tortuous but patent to the
limits of dissection. The main arteries and veins dissected are natural, bile staining of the intima is seen.

Respiratory system. The air passages contain bile stained muco-purulent material, dissection carried down to the smaller bronchi does not disclose any tumour-like lesions. The lungs are small, confluent broncho-pneumonia is seen in both lower lobes. Tumour growth is not visible in either organ nor in the pleurae.

Alimentary system. The oesophagus is normal. Old adhesions are present in the peritoneal sac, the greater omentum is inflamed and adherent to the undersurface of the liver. A partial gastrectomy has been performed. The pyloric stump is thickened due to diffuse increase of interstitial tissue, this ceases at the pylorus itself. A pedunculated growth 2 cm. diameter is present in the posterior wall of the antrum. The intestines show only putrefactive change. The liver, 1400 g., is externally normal, repeated close slicing fails to reveal any nodules. The gall-bladder and ducts are normal. (The pressure test is positive despite adhesions). The pancreas is much softened.

Genito-urinary system. The kidneys (left 210 g., right 160 g.) are normal. The ureters, bladder, and prostate gland are natural.

Central nervous system. The meninges are merely bile stained. The brain is normal externally and on dissection.

Endocrine system. (The hypophysis cerebri was over-looked). The thymus gland is atrophied almost to extinction. The thyroid gland is normal; both adrenal glands show only marked autolysis.

Microscopical examination:
General description of the lesions encountered.

This necropsy was performed two days after death and the preparations are consequently poor. Preservation of structure, i.e. microarchitecture, is reasonably good but the staining properties are badly impaired.

The lesions where present are recognisable as moderately late Hodgkin's disease. Fibrosis and necrosis are extensive and eosinophils cannot be identified. Whereas some latitude of appearance is discernible the growth is much the same wherever it occurs. Comparison with the original biopsy is difficult because of post-mortem change but in some fields the resemblance is close. In a few nodes the loss of lymphocytes is disproportionate and accentuates the impression that reticulum cells are unduly numerous.

Non-specific broncho-pneumonia is present; evidence of tuberculous disease is absent.

/Distribution
Distribution of the lesion.

Lymph nodes. There is striking persistence of non-affected lymph nodes; these are found in the neck, para-tracheal chains, right axilla, mediastimun, mesenteries, and left inguinal regions. In addition the antecubital and popliteal nodes are not diseased. The lesion is identified in the left axilla (four nodes), the para-aortic (retro-peritoneal) nodes, a chain along the right iliac vessels and the right inguinal region.

The spleen is extensively affected, haemosiderosis and necrosis are pronounced. Lesions are also confirmed in the liver. The femoral bone marrow shows almost complete gelatinous degeneration.

All other tissues examined proved negative. The gastric tumour is a papilloma.

Diagnosis and summary.

Atypical Hodgkin's disease, eosinophils lacking and aplastic anaemia. This case is very interesting because of the restricted distribution of the lesions. It appears probable that the severe anaemia determined death. Oedema is not a feature and lymphoid tissue survival is unusually great.
Mrs. Catherine Cameron, aet 31 years. Ref. P.M.155:52. G.R.I.

Anamnesis. Early in 1951 the patient complained of a painless swelling in the inguinal region. On 20th April 1951 an enlarged inguinal lymph node was removed for microscopical examination at Stracathro Hospital, Brechin. This was examined at Dundee Royal Infirmary and Hodgkin's disease was diagnosed, Ref. 1186:51 Dundee Royal Infirmary; Serial 217. Radiotherapy was instituted and ten treatments were given. The patient was pregnant at the time and later gave birth to a normal healthy baby. Thereafter she remained well till ten weeks ago when jaundice, anorexia, nausea, and vomiting supervened. She was admitted to Irvine Central Hospital and her condition improved sufficiently for her to return home after several weeks. Two weeks ago dyspnoea developed suddenly and she became very frightened and distressed. On 17th May 1952 she was admitted to the Gatehouse of Glasgow Royal Infirmary in a dying condition. She lapsed into coma and died 19th May 1952.

Report of Necropsy. 8 hours post-mortem.

External. A well built adult woman. There is slight anasarca of the lower limbs. Jaundice is not present.

Reticulo-endothelial system. Lymph nodes. The lymph nodes in the neck, axillae, and antecubital fossae are very greatly reduced in number. In the first site a very few were found after prolonged search; they are confined to the thoracic inlet. Nodes could not be found in either antecubital fossa nor in the right axilla. A solitary partly fat-replaced node was recovered from the left axilla.

The mediastinal lymph nodes are numerous and much enlarged. They are discrete, hard, and white. The largest measures 4 X 3 X 2 cm. The para-tracheal chains on both sides are similarly affected but the lesions cease abruptly at the root of the neck. Intra-pulmonary nodes are conspicuous in both lungs due to their participation in the morbid process.

In the abdomen lymphadenopathy is considerable. Those in the porta hepatis, the para-pancreatic group and the coeliac group are moderately enlarged and firm. The retro-peritoneal para-aortic group and those along the iliac arteries are hard and white but not much enlarged. One large node 3 X 2 X 2 cm. is producing distortion of the inferior vena cava at the level of the right renal artery. Several smaller nodes are pressing on the right ureter below this level. The nodes of both groins are enlarged, hard, and white on section, and on the left side further enlarged nodes are present in the lower angle of Scarpa's triangle. Popliteal nodes cannot be found.

Spleen. (420 g.) The gland is fairly uniformly enlarged but slightly /distorted
distorted due to nodules in its substance. The capsule is smooth and free from exudate. The cut surface reveals a pale pink homogeneous pulp in which are set scattered angular white suet-like deposits of Hodgkin tissue. These vary widely in size the largest being 2 cm. in diameter. The hilar lymph nodes are partly embedded in the splenic substance and present the same changes as the other abdominal nodes. Visible deposits of the new tissue cannot be seen in the liver.

Bone marrow. The red marrow of the sternum appears normal. In the thoracic and lumbar vertebrae several white nodules are seen. The upper third of the shaft of the right femur is occupied by soft red marrow which is homogeneous, tumour is not found. The marrow of the skull appears normal, it is very scanty. The nuclei pulposi of the spine are normal.

Respiratory system. The trachea is moderately congested and the lumen is occupied by whitish-yellow mucus and pus. The lower 3 cm. presents nodular tumour-like beading in the mucosa. These nodules are continued downwards into the main bronchi. They are firm, yellowish-white and 2-3 mm. diameter. Gross ulceration is not convincingly shewn at any level but some of the larger lesions are stippled with pink. Stenosis of the right main bronchus obtains. The left lung is well aerated and appears normal. The right lung is collapsed and several tumour-like craggy nodules are present below the pleural surface at the periphery. These measure up to 1 cm. diameter. On section the growths are white. A small (unmeasured) effusion of sero-fibrinous character is notable on the right side.

Cardiovascular system. The pericardial sac is distended by approximately 180 cc. of clear yellow fluid. Two small white tumour nodules approximately 1 cm. diameter are noted in the visceral layer just below the line of reflection from the aorta and pulmonary artery. Heart (275 g.): The organ is of normal size and configuration. The myocardium is of good quality and the chambers are natural. The aorta and main arteries are of small calibre, intimal change is limited to fatty streaking. Thrombosis is present in both femoral veins, the left vessel is completely occluded, but the right only partially.

Alimentary system. The oesophagus is natural. The peritoneum of the pelvis, especially that in the pouch of Douglas, is beset by numerous small nodules which appear to have developed deep to it and are thrusting forward into the membrane. Exudate is scanty. The stomach and intestines appear entirely healthy; the mesenteric nodes are small and inconspicuous.

The liver (1980 g.), is large. The shape, consistence, and colour are natural; repeated close slicing reveals only terminal congestion. The gall-bladder and ducts are normal. The pancreas is well preserved and is normal.

Genito-urinary system. The kidneys (left 150 g., right 170 g.) are normal /externally;
externally; there is some persistence of the foetal lobulation. Section reveals congestion but tumour deposit is absent. Slight hydronephrosis is appreciable on both sides but the pelves and ureters are clear of intrinsic growth. The para-aortic lymph node tumours are the apparent cause. The bladder is small and contracted. Several small aligned nodules are present in the trigone, one small polypoid outgrowth of firm white tissue is seen. The uterus and adnexa are natural.

**Endocrine system.** Pituitary gland. The gland is normal in size and nodules are not visible in the cut surface. Thyroid gland, small, otherwise natural. Adrenal glands. Both are normal naked-eye.

**Central nervous system.** The scalp, skull, and meninges are natural. The brain (1300 g.) is normal.

**Microscopical examination:**

**General description of the lesions encountered.**

The form of the lesion is highly cellular tumour-like Hodgkin's disease. Fibrosis is present in most sites, but is chiefly restricted to fibrils and fibrillae; large hyaline masses are rare. In several lesions morbid reticulum cell overgrowth is conspicuous but eosinophils are retained. Compared with the biopsy the growth is much more tumour-like. Microscopical evidence of tuberculosis is not found.

**Distribution of the lesion.**

Lymph nodes. Lesions are identified in nodes from the mediastinum, abdomen, and both groins. The single left axillary node recovered is unaffected but shews advanced fat atrophy. The disappearance of peripheral nodes including those of the cervical regions is unusually complete. In the bone marrow deposits of the new tissue are small, they are confirmed in the ribs and lumbar vertebrae but absent in the right femoral marrow. In the spleen the lesions are rather more cellular than usual but otherwise typical. The liver samples studied are normal; there is not even lymphocytic infiltration of the portal tracts. The adrenal glands and hypophysis cerebri were not sectioned. A nucleus pulposus examined shews no evidence of disease.

Extra-nodal lesions are confirmed in the trachea, bronchi, lungs, bladder, visceral pericardium, and pelvic peritoneum. In these sites there is destruction of the native tissues by the growth. Ectopic lymphoid tissue foci co-exist with the peritoneal lesions.

The kidneys, thyroid gland, pancreas, breast, and ovaries are normal.

**Diagnosis and summary.**

Cellular Hodgkin's disease shewing enhancement of tumour-like growth
and properties. The disease ran a relatively short course (known duration 15 months). There is definite failure to systematise throughout the reticulo-endothelial tissue but extra-nodal involvement is quite pronounced.

Anasarca, mild hydrothorax, and hydropericardium are notable, there is moderate survival of lymphoid tissue in para-enteronic sites.

Anamnesis. The patient was admitted on 3rd January 1953. She had been attending outpatient dispensary for one year and was being treated for myxosodena. In November 1952 she had begun to feel unwell and a temperature chart taken at home showed a morning elevation of temperature. About that time she noticed swellings in her neck, axillae and groins. In the following month she experienced back pains and sciatica. Biopsy was performed (Ref. G.R.I. 495:53, Serial 191). She received a course of nitrogen mustard with little result, and three doses of generalised radiotherapy with slight regression of the glandular swellings. Femoral thrombosis and dropsy supervened shortly before she died on 23rd March 1953.

Report of Necropsy.

11 hours post-mortem.

External. A well-built, well-nourished middle-aged woman. The skin of the right thigh is pigmented a delicate brown with darker freckles of the same colour. A small surgical scar in the left groin is similarly pigmented. There is moderate dropsy of both legs.

Reticulo-endothelial system. Lymph nodes. The superficial groups of cervical lymph nodes are small and inconspicuous on both sides, but the deeper groups, especially those along the carotid sheath are enlarged glaucous and fleshy. One node on the left side and lying on the vertebral column at the thoracic inlet measures 5 x 2 x 2 cm. On section all these nodes are pale pink, fibrosis is not evident. Nodes are not recoverable from the antecubital fossae. In both axillae several enlarged soft pinkish-white nodes are present, their number is few. The retro-sternal nodes of Stibbe are markedly increased in number and size and are indistinguishable from the anterior mediastinal groups which also shew a florid increase. In the mediastinum the size of the nodes is modest, most are below 2 cm. diameter, besides those mentioned above both para-tracheal chains are conspicuous. The remainder including the hilar nodes are small, anthracotic, and apparently clear of tumour. In the abdomen the mesenteric nodes are nearly all normal in number and appearance. The retro-peritoneal groups however, are enlarged, firm, and fleshy. There is a large ovoid tumour mass 24 x 15 x 10 cm. occupying the right flank which is apparently of lymph node origin (retroperitoneal). This is invading contiguous structures including the liver, right kidney, right adrenal gland and the right psoas muscle. There is a solitary enlarged node in the porta hepatis. On both sides the pelvic nodes are enlarged and fleshy. The groin lymph nodes are also increased in size, they are very soft. Lymph nodes are not seen in either popliteal fossa.

Spleen. 210 g. The gland is congested. Repeated close slicing reveals a single white tissue nodule 2 mm. diameter.

Bone marrow. The marrow in the sternum and lower vertebral bodies is /pink
pink and uniform, the bone is soft due to resorption. In the right femur, red bone marrow is confined to the upper half of the cavity, it is pale and uniform. Fat occupies the rest of the space.

**Cardiovascular system.** The pericardium shows some diffuse thickening at the base but tumour nodules are not visible in it. The heart (250 g.) shows moderate brown atrophy. Atheroma is remarkably slight throughout the arterial system. Recent thrombosis occludes the lower half of the inferior vena cava, where it is encircled by tumour, and both iliac veins and their tributaries are occluded.

**Respiratory system.** Pleurae. Left, normal. Right, extensive organising fibrinous exudate with loculated encystment of serous fluid is present. Lungs, left 660 g., right 600 g. An obsolete tuberculous primary complex is present in the left organ. The right lung is partly collapsed, there is palpable induration in the lower lobe. The bronchi and trachea show engorgement. On the right side the diaphragm is almost 2 cm. thick due to infiltrating white tumour spreading from below, the muscle fibres are visibly separated by the growth.

**Alimentary system.** A serous ascites (1000 ml.) is present. The oesophagus, stomach and intestines are normal. The liver (2330 g.) is uniformly enlarged, pallor is marked. Direct infiltration by tumour is seen in the undersurface; apart from this nodules are not present. The gall bladder and ducts are natural. The pancreas appears normal.

**Genito-urinary system.** Kidneys, (left 160 g., right unknown). The left gland appears normal. The upper pole of the right is directly infiltrated by soft white tumour, the cortex at this point is pale whitish pink and blurred-looking. The ureters and bladder appear normal. The uterus contains a single myoma 3 cm. diameter, the adnexa are normal.

**Endocrine system.** The hypophysis cerebri appears normal but is soft. The thyroid gland is uniformly enlarged, whilst retaining its contours it has been subtly replaced by soft pink tumour. It measures 6 X 5 X 3 cm. The cut surface shows several well demarcated nodules deep in the growth. Both adrenal glands are much destroyed by tumour, fragments of the right one are found with difficulty.

**Central nervous system.** The skull and meninges are normal. The brain 1350 g. is natural.

**Microscopical examination:**

*General description of the lesions encountered.*

Preservation of the material is regrettably poor and the illustrations are consequently indifferent. The majority of the deposits are more or less uniform reticulum cell sarcoma in which fibrosis is relatively slight. Reticulin formation is demonstrable. The type cell is a morbid...
mononuclear type similar to that seen in the biopsy specimen, eosinophils are absent. In several sites gross coarse fibrosis typical of late Hodgkin's disease is seen. In several nodes Hodgkin giant-cells abound. By virtue of the biopsy findings this case is interpreted as one of Hodgkin's sarcoma (Darthin's type II, reticulo-endothelioblastoma) but the neoplastic habit is enhanced to the point where the growth is actively aggressive and destructive. The metamorphosis is more complete than is usually seen, the resultant tumour being more atavistic and approaching the 'stem cell lymphoma' of Gall and Mallory. Tuberculous disease is confirmed in the lungs but the process is obsolete.

Distribution of the lesion.

All the cervical nodes examined shew the tumour, several contain large collagen deposits indicating previous Hodgkin's disease. The axillary nodes are similarly affected but one on the right side is only partly replaced. This node shews fat atrophy, the more atrophic portion is free of the growth. Stibbe's nodes and all the mediastinal ones are neoplastic. The retro-peritoneal and pelvic nodes are replaced by tumour, the former group merging with the large mass in the right flank. In both groins the tumour is identified in all but one node from the right side which is unaffected, it shews profound depletion of lymphocytes.

The tumour is present in the femoral bone marrow and it is extensively necrotic. One sample of marrow is free of growth and is hypocellular. The single splenic nodule is tumour in which central fibrosis is marked. Tumour is confirmed in the liver; in blocks taken at random well away from the deposits due to direct extension, the growth is not seen. Both adrenal glands are widely infiltrated, it is not possible to learn whether the growths are intrinsic but naked-eye their investment indicated invasion from without. The hypophysis cerebri is free of tumour.

The neoplasm is not present in a nucleus pulposus (lumbar region) examined nor in samples of omentum. Extra-nodal tumour is confirmed in the thyroid gland which is almost entirely destroyed. The large tumour in the right flank is the same neoplasm, destruction of skeletal muscle is extensive. Deposits of the growth are seen throughout the right kidney, the left gland is not involved. The diaphragm, pleura and right lung contain deposits of the tumour. Sections from the brain and optic nerves are normal.

Diagnosis and summary.

Hodgkin's disease and Hodgkin's sarcoma, with loss of pleiomorphism, the final picture approaching a pure reticulum cell sarcoma. In this case systemic involvement is a marked feature, the virtual replacement of the thyroid gland is interesting since this coincides with myxoedema.
diagnosed during life. Despite the relatively wide involvement one normal and one partly involved node survive. These findings are highly important since this case may well be regarded as "acute Hodgkin's disease". Cases of this type are generally believed to be truly systematised, (Callender, 1934). Anasarca, hydrothorax and ascites are notable. There is very little survival of normal lymphoid tissue.
Anamnesis. In May 1950 the patient was found to have pulmonary tuberculosis, mainly right sided. Collapse therapy was followed by a satisfactory response. Eventually in October 1951 he was declared cured and returned to work. In February 1953 he had a bout of persistent vomiting, and jaundice developed. He was admitted to Hairmyres Hospital the following month. On examination multiple glandular swellings were found and biopsy established the diagnosis of Hodgkin's disease (Ref. 804;53 Hairmyres Hospital, Serial 220). He was treated with blood transfusion, nitrogen mustard and trephine melanine with little benefit. He became progressively worse and was admitted to Glasgow Royal Infirmary at the end of June 1953. At this juncture he was extremely ill and despite blood transfusion his condition worsened rapidly till death on 5th July 1953.

Report of Necropsy. 20 hours post-mortem.

External. A tall cachectic young man; deep orange yellow jaundice is present. (weight of body = 134 lbs.)

Reticulo-endothelial system.

Lymph nodes: The cervical nodes are considerably enlarged and are unduly numerous. The superficial groups in the anterior and posterior triangles are hardest and their increase in size is the least. Their cut surfaces are generally evenly yellow but occasional white areas are seen. The retro-pharyngeal groups are thrown into unwonted prominence because of their gross increase in size, they are softer than the others. They form a columnar mass 10 X 3 X 2 cm., nodes being closely set mutually, but remaining discrete. Individual specimens measure up to 3 cm. diameter. The cut surfaces are all yellow with occasional petechiae.

The axillary nodes on both sides are small, hard, and few. Despite much search none are found in either antecubital fossa. A single small subcapsular node is present on the left side. The para-tracheal glands are numerous, enlarged, and rather fleshy. The retro-sternal nodes of Stibbe are readily seen due to their increase in size. They are dull red and soft.

The mediastinal lymph nodes are very numerous and moderately enlarged. In most cases they are pinkish-yellow and rather soft. The right inferior tracheo-bronchial node is incompletely replaced by pale yellow tumour-like tissue, a rind of dull red congested lymphoid tissue being visible. The pulmonary hilar nodes are small and slate colour on the left side, but are enlarged and firm on the right side.

The mesenteric nodes are small, soft, and generally inconspicuous, except in the lesser omentum and portal fissure, where enlarged fleshy
nodes are found. The para-pancreatic nodes are small, fleshy, and numerous. The para-aortic and pelvic glands are much enlarged, firm and increased in number. In the groins the glands are small, infrequent, and hard. The popliteal nodes are very small.

The spleen. (1000 g.) is uniformly enlarged and very numerous angular yellowish-white deposits of firm tissue project slightly from the surface. These deposits measure 3-4 mm. in diameter. The cut surface is evenly studded with them; the matrix is dull crimson.

Bone marrow. Uniform red marrow is visible in the bodies of the lumbar and thoracic vertebrae. The right femur contains red marrow in its upper third, below this level there is only fat. Tumour nodules are not seen. The bones of the skull also appear normal.

Cardio-vascular system. The pericardium is natural, approximately 20 cc. of bile-stained fluid occupy the sac. The heart 240 g. shows moderate atrophy. Atheroma is virtually absent from the aorta and main branches dissected.

Respiratory system. Lungs. Left 760 g. Oedema and congestion are notable, small lesions of obsolete tuberculosis are present in the apex. Right 600 g. Partial collapse and fibrous pleurisy are present. Active caseous foci, some with thick fibrous capsules, are seen throughout the upper lobe.

Alimentary system. Localised fibrinous exudate is seen over the viscera, especially the spleen and liver. Several gelatinous fibrin clots lie in the pelvis. Free fluid is small in amount. The stomach and intestines show only early putrefaction. The liver (2600 g.) is uniformly enlarged. Scanty whitish-yellow nodules, rarely above 2 mm. diameter are present under the capsule and throughout the viscous. The organ is rich reddish-brown. The gall-bladder is distended with orange bile, the pressure test is positive. The pancreas is natural apart from bile staining and softening.

Genito-urinary system. Kidneys, left 250 g., right 250 g. Apart from their large size they appear normal. The ureters, bladder, prostate gland, and testes, are normal.

Central nervous system. The brain 1460 g. is normal and unstained by bile although this pigment colours the meninges and choroid plexuses.

Endocrine system. The hypophysis cerebri is small and dull red. The thyroid gland is atrophic, leaf-like, and bilious. Only one parathyroid gland is found, a small one. The adrenal glands are small but otherwise normal.
Microscopical examination;
General description of the lesions encountered.

All the tumour deposits assume the form of Hodgkin's sarcoma (type II of Warthin, reticulo-endothelioblastoma). Morphology conforming to ordinary Hodgkin's disease is not encountered. The growth is a highly pleomorphic reticulum cell sarcoma with an high content of Hodgkin giant-cells, moderate fibrosis, and total absence of eosinophils. Histologically the growth exhibits low malignancy. An interesting feature which is fairly common in the lymph nodes is partial involvement of these structures.

Pulmonary tuberculosis is identified and acid/alcohol fast bacilli morphologically identical with myco. tuberculosis are demonstrable in the lesions.

Distribution of the lesion.

Lymph nodes. The lesion is identified in the cervical, right subcapsular, Stibbe's retrosternal chains, the axillary and mediastinal nodes. In the abdomen the retroperitoneal and some of the mesenteric nodes exhibit it. Further lesions of the same character are observed in the left groin nodes. Normal but atrophied nodes are found in the right and left groins and in the right popliteal fossa.

The femoral bone marrow contains small foci of the growth. In the spleen the tumour is seen together with haemosiderin deposits and caseating necrosis. The liver contains microscopical deposits in the portal tracts. The adrenal glands, hypophysis cerebri, a nucleus pulposus, and the omenta are negative.

The thyroid gland, samples of lungs and skeletal muscle are also free of the growth. The sections of brain and choroid plexus examined are natural.

Diagnosis and summary.

Hodgkin's sarcoma; in this case metamorphosis appears to be complete. There is failure to achieve general systematization. Dropsy is not manifest to any degree in this case and surviving lymphoid tissue is present in quantity.
Anamnesis. About June 1952 the patient complained of general malaise and a painless swelling in the left side of the neck. She was admitted to Hairmyres Hospital and biopsy of the cervical tumour established the diagnosis of Hodgkin's disease. (Ref. 1735:52. Hairmyres Hospital, Serial 219). A course of X-ray therapy was given, the exposure being local to the neck and thorax. General improvement with regression of the swelling ensued, but six months ago recurrence developed. Further irradiation failed to influence the disease and she became very weak. This progressed till she was bed-ridden, and she was admitted, dying, to Glasgow Royal Infirmary on 12th November 1953. At this time she had mild anaemia and a large left-sided pleural effusion. A single intravenous injection of nitrogen mustard was given, 5.0 mgm., without untoward effect but two days later she passed into the agony and died 16th November 1953.

Report of Necropsy. 26 hours post-mortem.

External. An extremely emaciated young woman. There is pallor of the integument but no jaundice. Glandular enlargements are not palpable. Weight of the body, 73 lbs.

Reticulo-endothelial system. 1. Lymph nodes. Superficial lymph nodes accessible to dissection are not present in the neck. A few deep ones are found at the root of the neck. They are small, firm, and ill-defined. The cut surfaces display shaggy outlines and a pinkish-white substance. Nodes are also recovered with difficulty and in small numbers from the axillae, chiefly on the left side. Antecubital nodes cannot be found. In the thorax, Stibbe's nodes are irregularly enlarged, particularly on the left side. The mediastinal groups are markedly enlarged and hard. Most are white with caseous foci centrally. The paratracheal chains are enlarged as far up as the thoracic inlet. The hilar and intra-pulmonary nodes are grossly enlarged and hard. On the left side several up to 4 cm. diameter are visible. They are continuous with diffuse massive growth replacing the lung substance of the upper lobe on this side.

The mesenteric nodes are small, scanty, and inconspicuous; on section they appear natural. Several retroperitoneal nodes in the para-aortic groups are hard and craggy but little enlarged. In the pelvis the nodes are small and very few. Both groins have very few nodes, one firm medium-sized one is seen on the right side.

The popliteal fossae are devoid of nodes.

2. Spleen. 60 g. The gland is small, pale, and soft. Repeated close slicing reveals always an homogeneous pale pink surface. Nodules are not present.

/3. Bone
3. **Bone marrow.** The skull is normal, the diploe is small. There is tumour-like infiltration of the periosteum of the internal aspect of the 8th left rib, the marrow cavity is small and dry. (Infiltration or fibrosis is visible in the adjacent intercostal muscles but this process is limited in extent). In the sternum and lumbar vertebral bodies the marrow is pale pink, nodules are not visible.

**Respiratory system.** The larynx, trachea, and bronchi are natural apart from purulent sputum in the latter. A serous effusion of 1200 cc. is present in the left pleural sac. The left lung is collapsed and most of the upper lobe is replaced by hard white or café-au-lait tinted tumour. In addition rounded nodules of similar growth are seen in the lower lobe. The right lung is less collapsed but contains numerous scattered tumour nodules 0.5 - 4 cm. diameter. The picture closely resembles bronchial carcinoma, but bronchial ulceration is not a feature. In the left parietal pleura over the 6th, 7th and 8th ribs, and interspaces, small beads of very hard pale cream tissue are seen.

**Cardiovascular system.** The heart is small and atrophic, dissection was not performed. In the pericardium three tumour nodules 4-5 cm. in their long axes are present in the parietal layer. The aorta is normal and free of atheroma.

**Alimentary system.** The peritoneum is healthy. The oesophagus, stomach and intestines shew only post-mortem change. The liver 1000 g. is small but normal to naked eye scrutiny, repeated close slicing does not disclose any focal lesions. The gall-bladder and ducts are natural. The pancreas is softened and discoloured.

**Genito-urinary system.** The kidneys, left 120 g., right 110 g. are natural, they are clear of tumour. The ureters and bladder are normal, likewise the uterus and adnexa. The breasts are pubertal.

**Central nervous system.** Brain, 1250 g. Normal.

**Endocrine system.** The hypophysis cerebri is small and red. The thyroid gland is small but otherwise normal. (Parathyroids, not dissected). The thymus is not recognisable as a separate structure, it is possibly taken up in the thoracic tumour. The left adrenal gland is partly enclosed by and possibly also invaded by hard white tumour. The right gland appears normal.

**Microscopical examination:**

**General description of the lesions encountered.**

The great majority of the lesions are recognizable as atypical Hodgkin's disease. It presents very similar morphology to the biopsy but eosinophils are particularly numerous in many sites. It is the diffuse fibroblastic type with swathes and whirls of fibroblastic cells dominating the picture. Hodgkin giant-cells are often distinctively rare, the appearances then recalling fibrosarcoma in general form but / suspiciously
suspiciously indolent. Focal deposits of the new tissue are quite common in nodes.

In a few sites morbid reticulum cell proliferation is notable but typical Hodgkin's sarcoma is absent.

In this case all the specimens taken for microscopy are poorly preserved and the staining properties are badly impaired. Microscopical evidence of tuberculosis is lacking.

**Distribution of the lesion.**

**Lymph nodes.** Lesions are confirmed in the cervical, axillary, para-tracheal and mediastinal nodes. In the first named group hyaline fibrosis is pronounced. Further lesions are seen in the retroperitoneal and groin nodes but not in the mesenteric ones. These latter are very small.

**Bone marrow.** The marrow is hypocellular, the sample is free from deposits of the new tissue.

**Spleen.** Simple atrophy is present, no other changes are visible. The liver is free from deposits. The left adrenal gland is enclosed by Hodgkin tissue and at one point there is penetration of the capsule, a zone of necrosis precedes the growing edge of the tumour. Deposits are not found in the gland itself. The systemic involvement is most impressive in the lungs, pericardium and periosteum of the 8th left rib. In the first named site caseous necrosis abounds but the cellular reaction of tuberculosis is not seen. The kidneys are normal post-mortem specimens.

**Diagnosis and summary.**

Atypical Hodgkin's disease; diffuse fibroblastic type. Naked-eye the morbid anatomy recalled neoplasm of lung (see illustration) strongly. There is failure to systematise and the maximum focus is thoracic. Apart from the lack of evidence of rapid growth the picture is tumour-like. Oedema is a feature in this case, it is chiefly expressed as hydrothorax. Survival of lymphoid tissue is moderate, the spleen being intact.
John Young, aet 58 years. Ref. P.M. 418:53. G.R.I.

Anamnesis. The patient was first observed to have a swelling in his neck by his physician; this was five or six years ago. This lump persisted without alteration in size and his general health was good up till 1952. In May of that year he developed pneumonia and pleurisy and was treated in Heathfield Hospital, Ayr, for eight weeks. He recovered fairly satisfactorily but a productive cough remained. He again developed pleurisy in June 1953 and was treated in Irvine Central Hospital for five weeks. Since then progressive tiredness, loss of energy and increasing shortness of breath have developed insidiously. Three months ago an irritating rash of ephemeral duration affected his face and arms. Just prior to admission to Glasgow Royal Infirmary anasarca developed. At this time generalised lymphadenopathy and splenomegaly were discovered. Biopsy (Ref. G.R.I. 4643:53, Serial 209) of a cervical node performed 13th November 1953 revealed highly cellular tumour-like Hodgkin's disease. The patient died five days later.

Report of Necropsy. 16 hours post-mortem.

External. A man of later middle-age. There is pronounced cachexia which is extensively concealed by dropsy except in the head. Jaundice is not present. (Weight of body = 103 lbs.)

Reticulo-endothelial system. Lymph nodes. Generalised lymph node enlargement is present. The nodes in the neck, axillae, mediastinum, extra-mesenteric abdominal groups, para-aortic region and groins are increased in number and size, the largest being the size of horse beans. All are rather soft, discrete and fleshy. The cut surfaces occasionally reveal fibrosis but are usually a homogeneous pinkish-grey. The mesenteric glands are not much increased in size. The nodes under the subclavious muscles and Stibbe's para-sternal nodes are prominent due to increase in size. A lymph node, 1 cm. diameter, is present in the subcutaneous tissue over the left scapula. Two very small fibrous nodes were found in the antecubital fossae, one on each side. Popliteal nodes could not be found after prolonged search.

Spleen. (230 g.). The viscus is uniformly enlarged. A recent superficial fibrinous exudate is notable on the surface. The viscus is soft and pliable, the cut surface is faintly congested and multiple small creamy white spots are scattered through it, it is equivocal whether they are deposits of the new tissue. (Liver). The portal tracts do not shew any visible deposits.

Bone marrow. The bone of the femur, sternum, lumbar spines and skull appears normal, there is little osteoporosis. In the femur red marrow is confined to the upper third. It is clear of visible tumour; in the lower two-thirds gelatinous degeneration is present. The sternal and rib marrow is red and also clear of tumour.
Thymus gland. The structure cannot be identified separately from the anterior mediastinal lymph nodes.

Respiratory system. The larynx and trachea are natural. There is moderate bronchiectasis of cylindrical type in the tubes to the lower lobes on both sides. The pleurae are thickened and present multiple dense adhesions to the lungs. Approximately 1,000 cc. of serous fluid are present in the thorax. Both lungs are heavy due to oedema and tumour. The upper lobes are congested and dropsical and clear of tuberculous or neoplastic disease. Tumour-like infiltration of greyish-pink colour is visible in the cut surfaces of the lower lobes. It is chiefly para-bronchial though isolated irregularly outlined nodules about 1-2 cm. are also seen. The large deposits are poorly demarcated. On palpation the affected tissue is craggy; some necrotic softening is present. The appearances suggest that the growth has emanated from the mediastinum.

Cardiovascular system. Pericardium. Several hard white nodules are visible in the inner aspect of the parietal layer at the base. They appear to be totally fibrous. Heart (300 g.). The myocardium is of good quality but rather chestnut coloured. The chambers are natural. Mitral valve. There is stenosis of button-hole type with rigid calcified margins; the orifice admits one finger. (The lack of right ventricular enlargement is notable). The remaining valves are natural, slight sclerosis of the cusps being the only feature of note. Moderate atheroma of nodular type affects the coronary arteries. The aorta is atheromatous but the lesions are mild and calcification is slight. Arteries. Mild atheroma is notable with advanced arteriosclerosis. The cerebral vessels show only the latter. Veins. Thrombosis is not present in the large vessels.

Alimentary system. Tongue. The lymphoid tissue at the root is copious, it does not appear involved in tumour. Oesophagus. Natural. The stomach and intestines do not shew any morbid features of note, in particular tuberculous ulceration is not seen. The liver (1,350 g.) is small, truly atrophic and shews terminal congestion. The vascular pattern is well demonstrated in the under surface due to atrophy of parenchyma. Tumour is not encountered on multiple close slicing. The gall-bladder and ducts are natural. Pancreas. Faintly orange-yellow in colour, otherwise normal.

Genito-urinary system. Kidneys (left 150 g., right 190 g.). The left is the seat of renal phthisis with cavitation and pelvic ulceration. Calcareous change is absent. Deposits of tumour cannot be found. The right gland presents two large depressed scars probably of ischaemic origin; a nodule of firm pearly-white tissue, 6 mm. diameter, is present in the pelvis. The gland is otherwise natural. The ureters are normal and the vesical reaches appear of the usual calibre. The bladder is healthy in appearance. The seminal vesicles contain partly calcified caseous
caseous debris and both epididymes are tuberculous. Both testes contain small hard white fibrous nodules in their substance.

Central nervous system. There is a small deposit of reddish-white material like old blood clot between the skull and dura at the torcular Herophili. The sinuses are healthy. The leptomeninges are dropsical. The brain (1,430 g.) is normal on dissection. (A few minute cysts are present in the left lateral lobe of the cerebellum near Deiters nucleus).

Endocrine system. The pituitary gland is soft but otherwise normal. The thyroid gland is small and the lobes are leaf-like. Two parathyroid glands were recovered, they appear normal. The adrenal glands are small but apparently free from disease.

Microscopical examination:

General description of the lesions encountered.

Although the material is only 16 hours post-mortem, the preservation is indifferent; in many of the larger lesions ante-mortem necrosis is pronounced.

The lesion presents the same appearance wherever it is found and is identical with the biopsy except where necrosis obtains. As in the nosology of the biopsy, it is equivocal whether it should be regarded as highly cellular tumour-like Hodgkin's disease or as reticulum cell sarcoma. Eosinophils are recognisable in the deposits and therefore the former category is probably more correct.

There is microscopical evidence of tuberculosis in the lungs, liver, spleen, epididymis, right renal pelvis and adrenal gland. Acid/alcohol fast bacilli identical with Myco. tuberculosis can be demonstrated only in the renal lesions.

Distribution of the lesion.

Lymph nodes. The lesion is present in the nodes of the neck, supraclavicular groups, axillae, para-tracheal chains, and mediastinum. The left suprascapular node which presented clinically is also involved. (A corresponding node on the right side does not exist.) The left ante-cubital fossa node is involved but the right gland is normal. In the abdomen the retroperitoneal nodes are involved likewise those in the pelvis and both groins. Due to accident the mesenteric nodes escaped examination.

Bone marrow. The lesion is not found in the marrow from the femur, a thoracic, and a lumbar vertebra. Minute tubercles are present in the femoral marrow.

Spleen. Minute lesions are identified together with caseous tubercles.
tubercles. In the liver only the latter are seen and they are rare.

The lesion is not found in the adrenal glands or hypophysis cerebri.

Extra-nodal deposits are restricted to the lungs and pericardium. Sections of the pancreas, cerebellum, testis, prostate gland, and epididymis are negative. In the kidneys and epididymis active tuberculosis is found. The nodule in the right kidney is a fibroma.

Diagnosis and summary.

Highly cellular Hodgkin's disease with co-existent tuberculosis of acute caseating type. There is widespread but not universal implication of lymphadenoid tissue. Anasarca and hydrothorax are present and there is pronounced reduction of normal lymphoid tissue.
Mrs. Margaret Royal, aet 72 years. Ref. P.M. 57:54. G.R.I.

Anamnesis. In March 1953 the patient developed a small swelling in the left side of her neck. This was resected after a provisional diagnosis of salivary calculus had been made. The biopsy shewed reticulum cell sarcoma, almost certainly of lymph node. (Ref. G.R.I. 1820:53, Serial 221). She received a course of radium therapy to the neck and has been attending the radiotherapy department at regular intervals since. In November 1953 she developed a cold with a productive cough and a swelling appeared in the right side of the neck. Since then she developed pneumonia and became bed-ridden. Dyspnoea became severe latterly. She died 11th February 1954.

Report of Necropsy. 4½ hours post-mortem.

External. A well-built somewhat obese elderly woman. Purpuric patches are notable over the limbs. There is oedema of the lower limbs. (Weight of body = 140 lbs.)

Reticulo-endothelial system. Lymph nodes. In the neck the superficial lymph nodes are small and inconspicuous. The few found appear scarred. The deeper nodes, particularly those above the thoracic inlet are considerably enlarged on both sides. They are soft and discrete. The cut surfaces are fleshy and pink with occasional haemorrhagic areas. Fibrosis is not apparent other than in the capsules. The supra-clavicular nodes on both sides are similarly affected. In the left axilla nodes are very scanty. A few nodes shew marked fat atrophy except in the case of one small gland which is white. The right axilla nodes are similar to those at the root of the neck; they are few in number. Nodes were not recovered from either antecubital region. The para-tracheal lymph nodes are normal in number but are individually enlarged up to 2 cm. in their long axes. They are soft, fleshy and pink. The cut surfaces shew extensive haemorrhage. The mediastinal group, particularly those in the anterior compartment, are grossly enlarged, being up to 4 cm. long. These shew anthracosis and considerable necrosis. The hilar and intrapulmonary nodes are considerably enlarged. They present the same features as the former group. The mesenteric nodes shew moderate increase in number and size. Those most affected are seen in the mesentery of the small intestine and in the lesser omentum. The retroperitoneal nodes, especially those about the aorta and its main branches are conspicuously enlarged. The nodes in the pelvis are large and very soft. Many are haemorrhagic. In the groins several large haemorrhagic nodes are present on both sides. A single small node is present in the right popliteal fossa. The left fossa does not contain nodes.

Spleen. (70 g.) The gland is small and atrophic; repeated slicing does not disclose any lesions.

/Bone
Bone marrow. There is resorption of cancellous bone in the ribs, sternum and vertebral column. The marrow is pale pink and homogeneous. In the right femur the upper two-thirds of the shaft is occupied by pale pink marrow, and fat. Nodules are not present.

Respiratory system. Trachea. In the lower third of the posterior aspect of the lumen several white nodules 2-3 mm. in diameter are visible growing in the mucosa. Further nodules are growing down into the bronchi on both sides. The growth is much more marked on the left side. It has brought about extreme stenosis of the left main bronchus. Lungs (left 370 g., right 490 g.). The left lung is almost entirely collapsed. The hilum is converted into a coarsely nodular pink and white tumour-like mass due to the grossly enlarged lymph nodes at this site. Several tumour deposits are present at the periphery of the organ; these measure 1-3 cm. in diameter. The right lung is fairly normal, apart from increase of nodules at the hilum and a single firm mass at the anterior lower margin of the lower lobe. This latter measures 1 X 2 cm. A serous effusion of approximately 300 ml. is present on both sides.

Cardiovascular system. Pericardium. The external layer is adherent to the enlarged glands of the mediastinum, but the membrane is not infiltrated. Heart (320 g.). Moderate hypertrophy of the right ventricle is the only morbid feature of note. Both coronary vessels are atheromatous. The aorta shows atheroma, with ulceration and calcification.

Alimentary system. The oesophagus is distorted by the mediastinal growth but is not infiltrated. The stomach is of normal size. The mucosa presents multiple small ulcers scattered throughout it. These have the appearance of malignant ulcers. The duodenum and small bowel are normal. The colon is natural. The liver (1,060 g.) is small. Dissection does not reveal any abnormalities. The gall-bladder and ducts are normal. The pancreas appears slightly scarred in places, but is otherwise natural.

Genito-urinary system. Kidneys (left 150 g., right 150 g.). The only abnormalities visible are several small white poorly demarcated nodules in the cortical tissue of each gland. The deposits are firm and measure 1-4 mm. diameter. The pelvis, ureters and bladder are healthy. The uterus contains a small endometrial polyp. There are several hyaline myomatous in the wall. Both ovaries appear natural.

Endocrine system. The pituitary gland appears normal. The thyroid gland (78 g.) shows rather diffuse enlargement of its left lobe. Definite tumour is not identified in it. The single parathyroid gland recovered is normal, but small. Both adrenal glands are small and not very well preserved; they are otherwise natural.

Central nervous system. The scalp, skull and meninges are natural. The brain (1,240 g.) is of normal size. Dissection reveals circumscribed gliosis in the right thalamus. This probably represents an old softening.
but orange staining is not visible. Recent softening is not visible. The vessels at the base are sclerotic and atheromatous.

**Microscopical examination:**

1. **General description of the lesions encountered.**

   In all sites where tumour is confirmed the type cell is a reticulum cell which shows some degree of lymphoid differentiation. The cytoplasm is feebly oxyphilic and indistinct. The nucleus is spheroidal, occasionally crenate or indented and confined by a sharply defined nuclear membrane. The nucleoplasm is not so leptochromatic as in the typical morbid reticulum cell but oxyphilic nucleoli are common. Reticulin and collagen formation are slight in most examples. In several sites eosinophils abound and the appearances approximate closely to cellular Hodgkin's disease. Compared with the biopsy the growth is more lymphoid in cell type.

   Microscopical evidence of tuberculosis is lacking, the only other pathological process is a minute chromophobe adenoma in the hypophysis cerebri.

2. **Distribution of the lesion.**

   **Lymph nodes.** The tumour is confirmed in the right cervical nodes, in one from the left side, in both supraclavicular groups and in the right axillary nodes. Both paratracheal chains, the mediastinal and intrapulmonary nodes all show the growth. Necrosis in these deep nodes is extensive. All the abdominal nodes taken show the disease. In both groins tumour is identified and is also present in a single node recovered from the right popliteal fossa. One left cervical node and one left axillary node are free from tumour.

   In the blocks taken tumour is not identified in the spleen, lumbar vertebral marrow, nor sternal marrow. Foci are present however in the femoral marrow. The liver is devoid of deposits likewise the adrenal glands and hypophysis cerebri.

   Extra-nodal deposits are confirmed in the trachea, bronchi, lungs, stomach, peritoneum and kidneys.

   The thyroid, parathyroids and pancreas are free of tumour.

**Diagnosis and summary.**

Reticulum cell sarcoma. There is evidence of instability of the microscopical picture. Systemic involvement is incomplete, several nodes, the liver and spleen not showing the lesion.

Anasarca and hydrothorax are present. Lymphoid tissue survival is somewhat greater than might be expected.
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1. Mesenchyme.
   Figs. 1-3
2. Lymph nodes.
   Figs. 4-36
3. Ectopic lymphoid tissue.
   Figs. 37-44
   Figs. 45-85
5. Hodgkin's sarcoma, Biopsy Series.
   Figs. 86, 87
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   Figs. 96-105
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   Figs. 106-109
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   Figs. 110-113
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11. Necropsy Series.
    Figs. 117-205
12. Peculiar fibrosis.
    Figs. 206-212
13. Lymphocytoid eosinophiles.
    Fig. 213
14. Experimental.
    Figs. 214-235
15. Lymph node counts.
    Figs. 236-246
Early mesenchyme. Note mutually similar trachychromatic nuclei with stellate cytoplasmic processes. Set in mucoid matrix. 6 mm Human embryo. H&E X 200

Fig. 1.

Nucleus pulposus. Large widely scattered spheroidal nuclei set in physaliphorous mucoid cytoplasm. The structure does not recall mesenchyme. Girl aet 13. H&E X 200

Fig. 2.
**Fig. 3.** Pure myxoma. The nuclei are closely set in this example, otherwise the resemblance to mesenchyme is near.
1332. H&E X 200.

**Fig. 4.** Axillary lymph node from a middle aged adult, the blood vascularity is prominent as a result of post mortem auto-injection.
P.M. 337:49. H&E X 150.
Axillary lymph node, from a woman aged 83 years. Depopulation of small lymphocytes is seen but sinus tissue persists and is not fibrous.

739. H&E X 90.

Same node as Fig. 5. High power view to show diminution of small lymphocytes, relative increase of reticulum cells and partial replacement by plasma cells.

739. H&E X 250.
LYMPHOCYTE DEPLETION IN NODE INVADED BY MUCOID CARCINOMA

Fig. 7  A groin lymph node invaded by mucoid carcinoma. A diffuse fibroblastic reaction is seen in the marginal sinus. Lymphocyte depletion is already well established.

3408:52  H&E X 175.
Fig. 8  Axillary lymph node showing moderate excavation by eccentric fat cone. Note capsule is thinned where fat reaches it (below) but does not collapse. Also note fine sprinkling of individual fat cells in lymphoid tissue beyond limit of fat cone. 311:50. H&E X 8.

Fig. 9  Axillary lymph node. Late fat replacement. Note lipoma-like uniformity of fat cone and early disappearance of capsule (right). 310:50. H&E X 11.
Fat replacement, high power view of edge of cone; note uniformity of fat cells and reduction of lymphocytes. The sinus at the junction resists the process.

3475:52. H&E X 200.

Fat replacement. Note persistence of sinus tissue which is outflanked by infiltrating fat.

3475:52. H&E X 200.
A peripheral solid follicle. Surviving sinus tissue is seen above. Hyalinised capillaries are well shewn, many exhibit stenosed lumina. The lymphoid tissue is clear of fibrosis.
Man 70 years. H&E X 250.

Advanced capillary hyalination. Lumina are absent and the collagen is increased by accretion. Note the marginal sinus is not fibrosed.
Woman 81 years. H&E X 270.
Fig. 14 Extreme vascular fibrosis. The line of marginal sinus can be traced, arrows. The vessels bottom left are possible new formations. Woman 73 years. H&E X 250.

Fig. 15 Late vascular fibrosis. Early calcareous change is present in the collagen, note the atrophy of the paratrabecular sinus tissue. Woman 75 years. H&E X 100.
Fig. 16 Vascular fibrosis of lymph node. An arteriole is seen at top left. Endarteritis is present and hyalinisation affects the pre-capillary branches. Sinus tissue persists.
Woman 61 years. H&E X 250.

Fig. 17 Para-vascular fibrosis of lymph node. The arteriole running in a trabecula surrounded by sinus tissue is appreciably thickened by concentric adventitial lamellae.
Man 50 years. H&E X 250.
Late diffuse fibrosis of the pulp. The deposit resembles scar tissue, note the thin walled vessels in it. (Para-vascular fibrosis is seen in the sinus below). This form of fibrosis is very rare.

Woman 57 years. H&E X 100.

Fig. 18 Para-vascular fibrosis of lymph node. Note how a mantle of sinus tissue persists round the axial fibrous tissue.

Man 62 years. H&E X 100.

Fig. 19
Diagrams 1-5 outline the development of lymph nodes.
Fig. 20 Early formation of lymphatic plexus at locus of developing lymph node.
22 mm Human Embryo. H&E X 200.

Fig. 21 Lymph node anlage, a small group of lymphocytes and reticulum cell nuclei are seen in the centre of fusing lymphatic channels.
46 mm Human Embryo. H&E X 250.
**Fig. 22** Lymph node showing central lymphoid tissue mass and marginal sinus. Note continuity with lymphatics.
65 mm Human Embryo. H&E X 125.

**Fig. 23** Lymph node showing very faint segmentation of central lymphoid mass. The marginal sinus is partly collapsed. At right an early trajectorial sinus is seen.
78 mm Human Embryo. H&E X 125.
Fig. 24  Early foetal lymph node. Sinus structure appearing, also possibly intra-nodal sinuses. Note afferent valved lymphatic at bottom right.
94 mm Human Foetus. H&E X 100.

Fig. 25  Foetal lymph node shewing approach of definitive blood supply. Small artery and vein.
108 mm Human Foetus. H&E X 125.
5^ months baby (premature)

Lymph node. The definitive blood supply is seen entering left and carries a mantle of sinus tissue round it.

5½ months baby. H&E X 90.

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Early segmentation of node substance. Elaboration of sinus tissue and early changes in lymphoid tissue density.

6 months baby (premature).

H&E X 90.
Fig. 28. Node at 7 months intra-uterine life. The sinus system is well developed, and a proper blood supply is present.
7 months baby (premature)
H&E X 90

Fig. 29. Same node as Fig. 28. High power to show detail of developing sinuses. The structure is already characteristic.
7 months baby (premature)
H&E X 200.
Fig. 30  Lymph node at birth. (Axilla). Several solid follicles are now present, and the node is properly formed. Child at term. H&E X 50.

Fig. 31  Adventitious lymphoid tissue formation outwith the capsule of a node. The vessel traversing the field obliquely delineates the capsule. Above it a follicle with a cuff of new sinus tissue has formed, outwith it is lymphoid tissue in direct contact with the adjacent areolar tissue. S.163:49. H&E X 200.
Neogenesis of lymph node in axillary fat. A lymphatic and capillary are related to focal lymphocytic aggregation. The deposit is not encapsulated.

Fig. 32 Neogenesis of lymph node in axillary fat. A lymphatic and capillary are related to focal lymphocytic aggregation. The deposit is not encapsulated.
S. 10:51. H&E X 150.

Fig. 33 Newly formed lymph node. Two sinusoidal capillaries and a lymphatic are seen in relation to the node which is acquiring a capsule.
Fig. 34  Same node as Fig. 33 at another level to show incipient pale centre follicle.  3.27:50.  H&E X 125.

Fig. 35  Minute newly formed node with sinus tissue.  From axillary fat.  147.  H&E X 200.
Fig. 36 Metastatic cancer in a newly formed node. It is sited in the node in a lymphatic, a true marginal sinus is absent. From mesentery of large bowel.
1462.    H&E X 65.

Fig. 37 Ectopic lymphoid tissue, prostate gland. Note pale centre of new follicle and displacement of blood vessel (right).
N. 51     H&E X 150.
Fig. 38  Ectopic lymphoid tissue in bladder. Solid follicle with scattered reticulum cells. 3727:50. H&E X 110.

Fig. 39  Ectopic lymphoid tissue in ureter wall, isolated follicle. Th.7. H&E X 200.
Fig. 40 Ectopic lymphoid tissue in female breast (chronic mastitis).
N.182. H&E X 250.

Fig. 41 Ectopic lymphoid tissue in gall-bladder.
3787:50 H&E X 100.
Fig. 42  Ectopic lymphoid tissue in skin (cutis vera) of face. Th. 3. H&E X 250.

Fig. 43  Small bronchus, showing morbid increase of peri-bronchial lymphoid tissue. Case of bronchiectasis. Unknown. H&E X 100.
Fig. 44  High power view of new lymphoid tissue to shew minute structure. (Bladder). 3727:50. H&E X 600.
HODGKIN'S DISEASE. Exodic growth phenomenon.

Fig. 45 Serial 70. Typical Hodgkin's disease showing exodic growth phenomenon, the dehiscent crescent is reactive lymphoid tissue, below is the new tissue.

H&E X 9.

Fig. 46 Serial 40. Atypical Hodgkin's disease. Exodic growth phenomenon visible but less pronounced. This lesion was tumour-like.

H&E X 7 ½
Folliculoid habit


**Fig. 49** Serial 56. Late Hodgkin's disease. Small deposits of active new tissue in a fibrous matrix.

H&E X 7½.

**Fig. 50** Serial 42. Early Hodgkin's disease. Reticulum cell proliferation with production of morbid forms. Note the typical 'owl-eye' nuclei.

H&E X 250.
Fig. 51 Serial 89. Early lesion. Pleomorphism of reticulum cells enhanced, very early fibrosis.
H&E X 250.

Fig. 52 Serial 89. Field shewing established lesion, fibrosis of fibril type is distinct.
H&E X 250.
HODGKIN'S DISEASE. Microscopical features.

Fig. 53 Serial 117. Relatively focal proliferation of reticulum cells; morbid mononuclear and epithelioid types are seen. These foci are quite common in Hodgkin lesions.
H&E X 250.

Fig. 54 Serial 184. Fibrosis in Hodgkin's disease. Fibrillae, fibrils and early coralline deposit are shown. Van Gieson X 110.
Fig. 55 Serial 83 (2nd biopsy). 
Late fibrosis in Hodgkin's disease. 
H&E X 250.

Fig. 56 Serial 184. Perivascular lamellae, artery and vein. 
H&E X 200.
HODGKIN'S DISEASE. Microscopical features.

Persistence of normal structure.

Serial 101. Necrotic focus in Hodgkin's disease. A thrombosed vessel is seen left.

H&E X 100.

Serial 10. Exotic growth phenomenon, ring of crushed lymphocytes and obliteration of sinus, the new tissue is sharply demarcated in this field.

H&E X 250.
Fig. 59  Serial 89. Surviving lymphoid follicle with pale centre, at periphery of node showing Hodgkin’s disease.

H&E X 200.

Fig. 60  Serial 199. Normal lymph cords and sinuses deep in a node showing Hodgkin’s disease.

H&E X 200.
Serial 77. Persistence of marginal sinus in atypical tumour-like Hodgkin's disease. This is rare.

H&E X 250.


H&E X 200.
HODGKIN'S DISEASE. Abnormal giant-cells.

**Fig. 63** Serial 153. Langhan's giant cell in Hodgkin's disease.
H& E X 200.

**Fig. 64** Serial 81. Hodgkin's disease in female breast. The new tissue in a gland field.
H& E X 200.
Fig. 65 Serial 81. Note breast acini in lower half of field. H&E X 250.

Fig. 66 Serial 206. Possible origin of Hodgkin's disease in the sinuses. Note morbid monocellular reticulum cells replacing littoral cells, the pulp is less affected. H&E X 200.

H&E X 250.
Figs. 69 and 70. Serial 153. Hodgkin's disease in which epithelioid cells are very numerous, symplasmic fibrosis is light; it is seen better in Fig. 70. 

H&E X 250.
ATYPICAL HODGKIN'S DISEASE. Cellular and tumour-like type.

Fig. 71 Serial 213. Highly cellular tumour-like Hodgkin's disease with retention of eosinophiles. Note relative uniformity of pattern. H&E X 200.

Fig. 72 Serial 204. Highly cellular and tumour-like eosinophiles reduced. Few very large cells are seen. H&E X 250.
ATYPICAL HODGKIN'S DISEASE. Cellular and tumour-like.

Fig. 73 Serial 31. Nosology doubtful, either cellular tumour-like Hodgkin's disease or reticulum cell sarcoma. Eosinophiles present. H&E X 250.

Fig. 74 Serial 209. Cellular tumour-like Hodgkin's disease, or reticulum cell sarcoma. Eosinophiles are very few. The picture is nearer frank tumour. H&E X 250.
Fig. 75  Serial 43. Reticulum cell sarcoma retaining many Hodgkin's disease features. H&E X 250.

Fig. 76  Serial 168. Diffuse fibroblastic Hodgkin's disease; pleomorphism limited, lymphocytes sparse, very uniform throughout the node. H&E X 250.
Fig. 77 Serial 44. Peculiar Hodgkin's disease. Remarkable overgrowth of morbid mononuclear and Hodgkin giant cells, fibrosis and eosinophilia present.

H&E X 90.

Fig. 78 Serial 44. Peculiar Hodgkin's disease. High power view of component cells. H&E X 600.
Serial 2. High power of Fig. 79 at capsule. Small lymphocytes predominate but several large cells are present.

H&E x 250.
Fig. 81  Serial 54. Benign Hodgkin's disease. (The preparation was poor). An Hodgkin giant-cell is seen at the right edge.

H&E X 250.
Figs. 82 and 83. Serial 23. A link lesion between ordinary and benign Hodgkin's disease. Fibrosis pronounced and small cells predominate.

H&E X 250.
Fig. 84  Hodgkin's disease with affinity to Brill's disease. Note follicular pattern with giant-cells in the false follicles.
1379:54.  H&E X 65.

Fig. 85  High power view of same lesion as Fig. 84. It is very close to Hodgkin's disease.
1379:54.  H&E X 250.
Serial 188. Hodgkin's sarcoma (Biopsy). Eosinophiles reduced but present, regarded as a transition stage.

H&E X 250.

Serial 202. Hodgkin's sarcoma (Biopsy specimen). Enhanced pleiomorphism, many giant cells, reduced lymphocytes, no eosinophiles, and fibrosis present.

H&E X 250.

Fig. 89. Serial 222. False Hodgkin's disease. Small islet of cancer cells which indicate the true nature of the lesion. H&E X 250.
Fig. 90 Serial 95. False Hodgkin's disease. Probable cancer metastasis. Diffuse fibrosis limited cellular pleomorphism. H&E X 250.

Fig. 91 Serial 95. False Hodgkin's disease. Plasmodial islet of epithelial cells which mimic morbid reticulum cells closely. H&E X 250.

Fig. 92 False Hodgkin's disease. Diffuse cancer metastases, from mammary primary (proved). H&E X 250.

Fig. 93
Fig. 94  Serial 86. High power view. Note that proliferated reticulum cells are small, even though nucleoli are common.

H&E X 600.

Fig. 95  False Hodgkin's disease. Proven tuberculosis of lymph node. The diffuse epithelioid cells mimic the lesion closely, but are small.

H&E X 200.

Serial 211. Area 'B'. Hodgkin's sarcoma. Large morbid reticulum cells very numerous. H&E X 250.
Fig. 98 Serial 188. Area 'A'
Hodgkin's disease. Fairly typical.
H&E X 200.

Fig. 99 Serial 188. Area 'B'
Transition features between Fig. 98 and Fig. 100.
H&E X 200.
Fig. Serial 188. Area 'C'.
100 Hodgkin's sarcoma.
H&E X 200.

Fig. Serial 22. Mixed lesion.
101 Ageing Hodgkin's disease area.
H&E X 250.
Fig. 102. Reticulum cell sarcoma area. H&E X 250.

Fig. 103. Coexistent Hodgkin's disease and reticulo-sarcoma in adjacent nodes, same biopsy. Hodgkin's disease node. H&E X 200.
Fig. Serial 191. Reticulum cell sarcoma node. (See also Fig. 103.)

H&E & 200.

Fig. Serial 85. Curious indeterminate lesion. ? Lymphoblastic Hodgkin's disease. Note very numerous large immature lymphocytes, reticulum cells and giant cells.

H&E X 200.
Fig. Serial 43. Reticulum cell sarcoma with affinities to cellular Hodgkin's disease. A morbid mononuclear reticulum cell is the type-cell.

H&E X 250.

Fig. Serial 82. Reticulum cell sarcoma, stem-cell type. Fibrosis tenuous. Mitoses numerous.

H&E X 250.
Fig. Serial 35. Reticulum cell sarcoma. Very large morbid mononuclear reticulum cells; some multinucleate; and fibrosis.

H&E X 250.

Fig. Serial 142. Reticulum cell sarcoma. Included normal lymph node in the tumour.

H&E X 7.
Serial 151. Lymphosarcoma, pure. Remarkable uniformity of tumour cells at this magnification.
H&E X 210.

Serial 151. H.P. field. Note the immaturity of the lymphocytes and the rare reticulum cells.
H&E X 600.
Serial 192. Lymphosarcoma. Rare morbid reticulum cells and Hodgkin type of fibrosis. H&E X 200.

Serial 192. Early lymphosarcoma showing exodic growth phenomenon. Note hyperplasia in the crushed rind tissue. H&E X 100.
Fig. Serial 108. (Re-cut). Brill's disease. False follicles very well seen.
H&E X 7.

Fig. Serial 108. Interfollicular fibrosis, fibrillae and fibrils shewing some compaction.
H&E X 250.
Serial 37. Indeterminate lesion, with affinities to Brill's disease. The follicles are very large.

H&E X 5.
Fig. 117. A field from the edge of an axillary lymph node showing late Hodgkin's disease. Necropsy 'A' H&E X 200

Fig. 118. Mediastinal lymph node. Hodgkin's disease area. Necropsy 'A' H&E X 200
Lymph node from groin. Established Hodgkin's sarcoma. There are many pleiomorphic reticulum cells, many of them multinucleate, set in a dense collagen fibre mesh.

Necropsy 'A'  H & E X 250

Cervical node portion showing Hodgkin's sarcoma.

Necropsy 'A'  H & E X 250
Fig. Right cervical node. Note the whorling of fibroblastic cells. Necropsy 'A' H & E X 110

Fig. Node from upper border of pancreas. The tumour consists chiefly of large morbid mononuclear reticulum cells, multinucleate forms are less numerous. The collagen mesh is tenuous. Necropsy 'A' H & E X 250
Fig. Rib. Deposit of Hodgkin's sarcoma. Necropsy 'A' H & E X 100

Fig. Liver. Deposit of Hodgkin's sarcoma. A tumour cell can be seen in a sinusoid at the margin of the normal tissue. Necropsy 'A' H & E X 250
Fig. 125. Multinucleate giant-cells from deposit of Hodgkin's sarcoma in the liver. Note the densely staining nucleoli. In the largest cell ingested neutrophils are visible in the cytoplasm.
Necropsy 'A'  H & E  X  750

Fig. 126. Prostate gland, deposit of Hodgkin's sarcoma.
Necropsy 'A'  H & E  X  110
Fig. Biopsy. Highly cellular pleiomorphic Hodgkin's disease shewing well developed fibrosis. Serial 188 H & E X 200

Fig. Biopsy. From the same node. An area in which reticulum cell overgrowth is unusually florid. Serial 188 H & E X 200
Fig. 129 Necropsy. Spleen. In the lower half of the field Hodgkin's disease is present; in the upper half, Langhan's giant cells are seen at the periphery of tubercle follicles. In the upper right corner caseous material is visible.
Necropsy 'B' H & E X 125

Fig. 130 Necropsy. Transverse section of right ureter shewing a deposit of Hodgkin's disease in the wall. Note the transitional epithelium and dilated vessels.
Necropsy 'B' H & E X 125
Necropsy. Deposit of Hodgkin's disease in the lung, note the small areas of caseous necrosis.

Necropsy 'B' H & E X 125

Necropsy 'B' H&E X 250
Fig. Necropsy. Tuberculous follicles in the liver.
Necropsy 'B' H&E X 125
Original biopsy, 1946.
Cervical lymph node. The reticulum cell proliferation is marked, in addition to the usual morbid mononuclear and Hodgkin giant-cells, epithelialoid cell islets are visible. The nuclei of the latter are relatively feebly stained. Note the large slightly immature lymphocytes.
Serial 117. H&E X 250

Necropsy. Right inferior tracheo-bronchial node. Maturing Hodgkin's disease, lymphocytes are scanty and fibrosis is marked. The lesion is becoming distinctly tumour-like.
Necropsy 'C' H&E X 250
Fig. 136. Necropsy. Juxta-splenic lymph node. Large morbid reticulum cells are much more abundant, the lesion approximates closely to reticulum cell sarcoma. (Hodgkin's sarcoma).
Necropsy 'C' H&E X 250

Fig. 137. Necropsy. Tumour of stomach. In this site the tumour is frankly sarcomatous. The cells are mutually similar but some nuclear pleiomorphism persists. In the lower part of the photograph the pale tissue is smooth muscle being infiltrated and overrun by the growth.
Necropsy 'C' H&E X
Biopsy. Early Hodgkin's disease. Reticulum cell proliferation is evident but fibrosis is scanty and small lymphocytes are still very numerous.

Serial 72  H&E X 250

Necropsy. Left cervical lymph node I. Cellular picture of Hodgkin's disease but note scanty fibrosis and survival of marginal sinus on left of plate. The lymphocytes are large.

Necropsy 'D'  H&E X 250
Necropsy. Para-tracheal lymph node II. Peculiar giant-cells. These are similar in some respects to the Warthin-Finkeldy giant-cells or "inverted" Touton type cells.
Necropsy 'D'  H&E X 600

Fig. 141

Necropsy. Left cervical lymph node II. Overgrowth of large morbid mononuclear reticulum cells and marked reduction of lymphocytes.
Necropsy 'D'  H&E X 250

Fig. 140
Necropsy. Left groin node. Reticulum cell sarcoma. Uniform cellular tumour showing incipient fibrosis.
Necropsy 'D'  H&E X 250

Necropsy. Liver. H.II. Reticulum cell sarcoma. Note the uniformity and dense staining of the nuclei.
Necropsy 'D'  H&E X 250
Fig. 144. Necropsy. Left axillary lymph node showing fat atrophy. Note the follicles and normal marginal sinus.
Necropsy 'D'. H&E X 10
Biopsy. Cervical node, anatomical medulla. Clusters of morbid reticulum cells are set in a matrix of lymphocytes and delicate strands of collagen.
Serial 57. H&E X 250

Biopsy. The edge of the cervical node. On the left the capsule is visible adjacent to a zone of crushed lymphoid tissue. The right side of the field contains the new growth with very numerous aggregated morbid reticulum cells among which mitoses are frequent.
Serial 57. H&E X 250
Necropsy. Para-aortic retroperitoneal lymph node.
Sclerosing Hodgkin's disease with numerous morbid reticulum cells. Most of this node presents this appearance.

Necropsy 'E'  H&E X 250
Necropsy. Mesenteric lymph node 'B'. Transition stage of the lesion. It is equivocal whether this is the ordinary or sarcomatous form. Necropsy 'E'  H&E X 250

Necropsy. Edge of a deposit in the lung. The tumour occupies the alveoli and destroys the alveolar walls.
Necropsy 'E'  H&E X 250

Necropsy. Trachea. The ciliated columnar epithelial lining is visible in the upper edge of the field. The submucosa is occupied by tumour which has destroyed most of the native tissue.
Necropsy 'E'  H&E X 150
Necropsy. Solitary follicles of normal lymphoid tissue in the colonic submucosa.

Necropsy 'E'  
H&E X 250

Necropsy 'F' H&E X 200

Fig. 154

Necropsy. Cervical lymph node. Late typical Hodgkin's disease.

Necropsy 'F' H&E X 250

Fig. 155
Necropsy. Infiltration of myocardium by Hodgkin's disease, note the disruption of the muscle fibres and fat cells in the new tissue.

Necropsy 'F' H&E X 100

Necropsy. Peri-adrenal fat over-run by Hodgkin's tissue. The adrenal gland is intact.

Necropsy 'F' H&E X 90
Fig. Necropsy. Inguinal lymph node unaffected by the disease.
Necropsy 'F' H&E X 12
Fig. Biopsy. Right inguinal node; established Hodgkin's disease. Note the pleiomorphic reticulum cells, some of which are necrotic. The homogeneous material at bottom left is caseous.
Serial 218  H&E X 250

Fig. Necropsy. Right inguinal node; Maturing Hodgkin's disease. Note fine diffuse fibrosis and paucity of lymphocytes. Necropsy 'G'  H&E X 250
Necropsy. Para-aortic lymph node. Hodgkin's disease. Reticulum cells are numerous and lymphocytes scanty, necrosis is also a feature.

Necropsy 'G'  H&E X 250
Necropsy. Mesenteric lymph node. The capsule is at the top of the plate. Sinus catarrh of dropsical type is seen. The lymphoid tissue is hypocellular but clear of the disease.
Necropsy 'G' H&E X 250

Necropsy. Femoral bone marrow. The aggregate of pyknotic nuclei represents the largest survival islet of haemopoietic tissue in this section. The gelatinous material appears fibrillary due to artefact.
Necropsy 'G' H&E X 250

Necropsy. Mediastinal node. Diffuse fibroblastic transformation prominent, but pleomorphism is maintained.

Serial 217 H&E X 250
Fig. 167 Necropsy. Retroperitoneal node. Highly pleomorphic cellular Hodgkin's disease. Eosinophils are retained. Necropsy 'H' H&E X 250

Fig. 168 Necropsy. Juxta-pancreatic node. Particularly cellular lesion, fibrosis is less in evidence. Necropsy 'H' H&E X 250
Fig. 169. Necropsy. Rib. (Decalcified in Susa fixative). Typical Hodgkin's disease deposit. Necropsy 'E' H&E X 250

Fig. 170. Necropsy. Bronchus. Several mucin glands are seen in the lower part of the field, embedded in Hodgkin tissue. Necropsy 'H' H&E X 250
Fig. 171  Necropsy. Bladder. Sub-mucosal deposit of Hodgkin's disease, the overlying mucosa is lost. 
Necropsy 'H'  H&E X 10

Fig. 172  Necropsy. Base of tongue. Deposits of normal lymphoid tissue. Germinal centres can just be made out in two of the foci. 
Necropsy 'H'  H&E X 6½
Fig. Necropsy. Left axillary lymph node. Advanced fat atrophy is present. The tissue is otherwise physiological. (The section is imperfect due to artefact).
Necropsy 'H' X 10
Fig. 174. Biopsy. Field from one of the larger groin nodes; typical Hodgkin's disease.
Serial 191. H&E X 200

Fig. 175. Biopsy. Field from the smallest groin node; reticulum cell sarcoma.
Serial 191. H&E X 200
Fig. Necropsy. Cervical lymph node. Area of mature Hodgkin's disease. Note the proliferating reticulum cell nuclei at the margin of the collagen, lymphocytes are in abeyance. Necropsy 'I'  H&E X 250

Fig. Necropsy. Mediastinal lymph node. Polymorphic reticulum cell sarcoma, several Hodgkin giant-cells are present. Necropsy 'I'  H&E X 250
Necropsy. Psoas tumour. Partly necrotic skeletal muscle fibres are seen embedded in the tumour cells. Necropsy 'I' H&E X 250

Necropsy. Kidney. A surviving glomerulus is shown set in a matrix of tumour. Necropsy 'I' H&E X 250
Fig. Necropsy. Lymph node from right groin. The structure is not morbid beyond showing extreme lymphocyte depletion. Necropsy 'I' H&E X 250
Biopsy. Cervical lymph node. Fairly cellular Hodgkin's disease, a swathe of young connective tissue is seen in the upper part of the plate.
Serial 220. H&E X 200

Necropsy. Right inferior tracheo-bronchial node. A random field of the tumour now present. (All the lesions present this appearance). Hodgkin's sarcoma, note the numerous Hodgkin giant-cells, sclerosis and general pleiomorphism.
Necropsy 'J' H&E X 250
Necropsy. Lymph node from right popliteal fossa. This node is physiological. An high power view to shew normal marginal and trajectorial sinuses together with lymphatic cords. The capsule is seen, top right.

Necropsy 'J'  H&E X 250
Fig. 185 Necropsy. Lung; right upper lobe. Acute exudative and caseating tuberculosis. (Mycobacterium tuberculosis is demonstrable with Ziehl-Neelsen stains). Necropsy 'J' H&E X 250
Biopsy. Cervical lymph node. The cellular new tissue is disposed in irregular folliculoid or segmented fashion with paler staining connective tissue between the foci.
Serial 219  H&E X 6

Fig. 186

Biopsy. Cervical node. Diffuse fibroblastic Hodgkin's disease. Swathes of young connective tissue are seen arranged in a sinuous manner. Note the relative homogeneity of the growth.
Serial 219  H&E X 150

Fig. 187
Biopsy. A high power view of the edge of one of the nodules. Morbid mononuclear and Hodgkin giant-cells can be made out. The small deeply staining binucleate cells are eosinophils.

Serial 219    H&E X 250
Fig. 189. Necropsy. Cervical and thoracic viscera, Anterior Aspect. Note large tumour-like mass occupying position of thymus gland, and nodular masses in the pericardial sac. Intra-pulmonary nodules are just discernible. Necropsy 'K' $\frac{1}{2}$ Natural Size.
Fig. 109 B. Necropsy.
Posterior view of cervical and thoracic viscera; a sagittal slice has been made. Note the remarkable resemblance to carcinoma.
Necropsy 'K'  X\frac{3}{4}  Natural Size.
Necropsy. Mediastinal lymph node showing focal Hodgkin's disease. The pale nodules are the morbid tissue. Necropsy 'K' H&E X 8
Necropsy. Mediastinal lymph node. Diffuse fibroblastic Hodgkin's disease. This field is typical of most sites, there is general failure to form dense deposits of collagen.
Fig. 191 Necropsy 'K' H&E X 250

Necropsy. A cellular focus from the same node pictured in Fig. 191. These foci are small and rare, eosinophils are present in them.
Fig. 192 Necropsy 'K' H&E X 250
Fig. 193  Necropsy. Mediastinal lymph node. Same node as Fig. 191. Above and to the left cellular Hodgkin's disease is seen, below and to the right is normal reactive lymphoid tissue. An artefact cleavage space partly separates the two.
Necropsy 'K'  H&E X 250

Fig. 194  Necropsy. Cervical lymph node. Persistent cellular focus.
Necropsy 'K'  H&E X 250


Necropsy. Axillary lymph node. Preservation is not so good as in the biopsy specimen but the growth is essentially similar. Necropsy 'L' E&E X 250
Necropsy. Antecubital lymph node. The same lesion is present as in figs. 197 and 198.
Necropsy 'L' H&E X 250
Necropsy. Periphery of a nodule in the lung. The new tissue is diffuse and less coherent than usual due to post-mortem change.
Necropsy 'L'  H&E X 250

Necropsy. Liver. Indeterminate lesion; probably tuberculous. Most of the cells and nuclei in the focus are small. Early necrosis is present.
Necropsy 'L'  H&E X 250
Biopsy. General appearance of the tumour, a fairly uniform medium-sized morbid reticulum cell growth. Fibre formation is slight. Serial 221. H&E X 250

Necropsy. Right cervical lymph node replaced by tumour. The growth is symplasmic in the field shewn but most examples closely correspond to the free cell variety seen in the biopsy. Necropsy 'M' H&E X 250
Necropsy. Left cervical lymph node. Fat atrophy was notable in the node, the surviving rind tissue seen here is normal apart from diminution of lymphocytes. Note blood vessels and surviving lymph sinuses. The capsule is seen at top right.

Necropsy 'M' H&E X 250
Fig. Low power view of cheloid, following surgical wound. Note indistinct encapsulation. H&E X 5.

Fig. High power view of normal junction. H&E X 200.
Fig. Desmoid tumour, skeletal muscle giant cells surrounded by tumour tissue. 
H&E X 200.

Fig. Desmatofibroma. The collagen mesh recalls the filigree of Hodgkin's disease. 
H&E X 200.
Fig. Desmatofibroma, general view showing giant cells. H&E X 200.

Fig. Reidel's struma. Skeletal muscle overwhelmed by periglandular fibrous tissue growth. H&E X 100.
Fig. Myelofibrosis. Uniform young connective tissue replaces the marrow.

H&E X 175.

Fig. Lymphocytoid eosinophiles in loose lymphatic tissue of a lymph node shewing early Hodgkin's disease.

Dominici X 600.
Liver and spleen of rat L.3 in section. After weekly I.P.I. 10 mgm. trypan blue over 125 days.

Fig. Liver and spleen of rat D.3. in section. After weekly I.P.I. 10 mgm. trypan blue over 125 days. ½.

Fig. Left. Liver of rat A.5, after fortnightly I.P.I. 10 mgm. trypan blue over 330 days. Tumour-like nodules are not present. Right. Liver of rat A.4, after fortnightly I.P.I. 10 mgm. trypan blue over 200 days. Numerous spheroidal tumour-like nodules. 1/1
Fig. 217 Rat A.2. Trypan blue I.P.I. 182 days. Liver. Nodular deposits of cells along portal tracts. The dye-laden cells appear black. H&E X 65

Fig. 218 Rat A.2. Liver. HP view of previous figure. Note uniformity of cell type except for dye and iron holding cells. H&E X 250
Hat A. Liver tumour 217 days. Growth becoming diffuse and infiltrating liver tissue. H&E X 250.

Fig. 219

Hat 3. Liver tumour. 360 days. Exposure to dye ceased at 120 days. H&E X 250.

Fig. 220

Fig. Rat A. Liver tumour. 217 days. Growth becoming diffuse and infiltrating liver tissue. H&E X 250.

Fig. 219

Fig. Rat E. Liver tumour. 360 days. Exposure to dye ceased at 120 days. H&E X 250.
Fig. 221. Rat E.9. Liver tumour, silver impregnation to show reticulin formation. Silver X 250.

Fig. 222. Rat A.4. Liver tumour. Incipient fibrosis. H&E X 250.
Rat A.7. Liver tumour with central necrosis (280 days). H&E X 65.

Rat A.2. Spleen with capsule and omental tag above. Reactive hyperplasia and megakaryocytes. Omental tag shows similar picture to hepatic tumour. H&E X 250.

Rat 3.7. 252 days. Axillary lymph node showing light fibrosis, an isolated finding. The deep sinus tissue survives. H&E X 200.
Fig. Rat A.2. Lung. Perivascular and peri-bronchiolar cuffing with macrophages. H&E X 65.

Fig. Rat A.2. Kidney. Diffuse macrophage focus, simulates pyelonephritis. H&E X 60.
Mouse F.5. 1.0 mgm. trypan blue I.P.I. weekly. 250 days. Liver. Doubtful nodule in portal tract. Kupffer cells prominent due to dye content. H&E X 250.

Fig. 231. Mouse F.5. Spleen. Red pulp hyperplasia and mild haemosiderosis. Cells with dark cytoplasm contain iron. 
H&E X 250.

Fig. 232. Mouse F.20. Lymph node. Mild non-specific reaction. 
H&E X 250.

Rat C. 8. Vital new red. 10 mgm. I.P.I. fortnightly 360 days. This is the largest periporal lesion found in the series. HaE X 250.
Fig. 235. Rat C.8. Spleen, to show haemosiderosis. Prussian blue & saffranin. Red filter X 250.
Photographs Fig. 236 to Fig. 245 illustrate the lymph nodes recovered by dissection from cases 2, 3, 5, 6, 8, 9, 10, 11, 14, 20. Fig. 246 depicts nodes recovered from the left axilla in an untreated case of Hodgkin's disease; 46 masses are shown but if the lobulated nodes are considered fusion products the total is 55.

Text reference p. 109 et seq.

**Fig. 236**

**E.149.**
LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF AN INFANT AGED 6 DAYS. TOTAL NUMBER 10.

**Fig. 237**

**B.147.**
LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF AN INFANT AGED 30 DAYS. TOTAL NUMBER 23.

**Fig. 238**

**150.**
LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF A BOY AGED 5 YEARS. TOTAL NUMBER 32.

**Fig. 239**

**410.**
LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF A GIRL AGED 13 YEARS. TOTAL NUMBER 31.
LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF A WOMAN AGED 22 YEARS.
TOTAL NUMBER 26

LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF A MAN AGED 29 YEARS.
TOTAL NUMBER 29.

LYMPH NODES RECOVERED FROM THE LEFT AXILLA OF A WOMAN AGED 47 YEARS.
TOTAL NUMBER 24.
LYMPH NODES RECOVERED FROM THE
LEFT AXILLA OF A MAN AGED 61 YEARS
TOTAL 12.

LYMPH NODES RECOVERED FROM
THE LEFT AXILLA OF A WOMAN
AGED 81 YEARS. TOTAL 11.

Fig. 244

Fig. 245

Fig. 246
VOLUME IV

DRAWINGS, DIAGRAMS AND GRAPHS
# DIAGRAMS AND DRAWINGS

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MAXIMOW' SCHEMA OF LYMPH NODE (AND SPLEEN) CELL GENEALOGY, SLIGHTLY MODIFIED.
KEY.

\[\text{\(\uparrow\)}\] = Line of development with potencies unrestricted.

\[\text{\(\downarrow\)}\] = Line of development with potencies restricted.

\[\text{2}\] = Haemocytoblasts.

\[\text{3}\] = Free histiocytes = histiogenous wandering cells.

\[\text{4}\] = Monocytes.

\[\text{6}\] = Secondary erythrocyte series (Definitive).

\[\text{7}\] = Granulocytes.

\[\text{0}\] = Small lymphocytes.

\[\text{A}\] = Common fixed mesenchymal cell.

\[\text{AA}\] = Cell lineage of fixed syncitial histiocytic elements in the lymph nodes.

\[\text{BB}\] = Cell lineage of free independently proliferating haemocytoblasts in the lymph nodes.

\[\text{CC}\] = Cell lineage of fibroblasts in the lymph nodes.

NOTES.

The large lymphocyte of Maximow is identified with the lymphoid wandering cell and the haemocytoblast, (of Ferrata et alii).

There is not any indication of megakaryocytes being formed in lymph nodes, i.e. from haemocytoblasts, nor is there any reference to mast cell origin. The polyblast of Maximow is conveniently regarded as an "indeterminate" cell which has several possible origins, and several possible development lines, and thus represents a kind of neutral phase.

In extra embryonic life, fibrosis can arise from reticular syncytium, both directly and via a polyblast metamorphosis.
LYMPH NODES AND SPLEEN.

FIBROBLASTS. RETICULAR HAEMOCYTOBLASTS.
SYNCYTIUM.

EXTRA-EMBRYONIC.

EMBRYONIC.

COMMON FIXED MESENCHYMAL CELLS.

FREE HISTIOCYTES.

FIXED HISTIOCYTES.

FIBROBLASTS.

ENDOTHELIUM.
GENERAL RELATIONS OF THE LYMPHATIC SYSTEM

Arterial distribution is in red, the venous in blue. The vertical blue/red line represents the blood capillary mesh. The lymph stream is shown in yellow.

Most lymphocytes entering the lymph stream are derived from the para-enteronic lymphoid tissue and related lymph nodes. The main lymphatic duct effects junction with the termination of the venous system. (The lymphatic drainage from the lungs is omitted in this diagram).
GENERAL RELATIONS OF THE LYMPHATIC SYSTEM

RA
LA
RV
LV
LYMPHATIC PLEXUS.
LYMPH NODE.
LYMPH NODE.
MESENTERIC LYMPH NODE.
LUMEN OF G. I. TRACT.
**SCHEMATIC LYMPH NODE**

A. Afferent lymphatic piercing capsule to reach marginal sinus.

A'. Efferent lymphatic leaving in hilar intrusion.

B. Capsule.

C. Marginal (sub-capsular) lymphatic sinus.

D. Cortical germinal lymphoid follicle, active.

E. Inactive germinal follicle.

F. Solid (secondary) lymphoid follicle.

G. Trabecula in section, note cuff of trajectorial lymph sinus.

H. Pulp of lymphoid tissue (loose lymphatic tissue, functional medulla).

I. Vessels of hilar intrusion.

J. Medullary lymphatic sinuses.
Three hypothetical sections differently treated.

Fig. i Silver impregnation.

Fig. ii Ideal pencilled preparation.

Fig. iii Routine haematoxylin and eosin stained paraffin section.

In Fig. i. the argyrophile fibres - reticulin is shewn, note the absence from the pale centre. In Fig. ii. the reticulum cell plasmodium is shewn, pervasive continuity is maintained. Fig. iii depicts the familiar appearance of lymph node sections.
FAT REPLACEMENT OF LYMPH NODES

Figs. 1 and 2 represent low-power and high-power views of the process.

In Fig. 1. Note that the fat of the fat cone is composed of larger cells than the fat of the adnate external areolar tissue. The lymphoid tissue has fallen back onto the persisting capsule and some marginal sinus tissue persists. Below the fat cone has reached the capsule and this membrane is disintegrating.

In Fig. 2. Note the attenuation and complete lack of reaction of the lymphoid tissue, at this point the tissue resembles marrow. Lymphocytes tend to persist between the fat cells.
FAT REPLACEMENT OF LYMPHOID TISSUE IN A LYMPH NODE

This high power drawing illustrates an interesting point. The node was recovered from an elderly man who had been tattooed when he was a soldier fifty-two years previously. Residual granules of insoluble dye-stuff are seen in the cytoplasm of macrophages at A, A' and in a fat-cell cytoplasm at A 1. Note the minute epinuclear vacuole in the nucleus of A 1 - this is seen only in fat cells. The finding suggests that a process of metamorphosis from reticulum cell to fat cell is part of the mechanism of fat atrophy. (The drawing is from the edge of the fat cone).
VASCULAR HYALINISATION FIBROSIS IN A LYMPH NODE

A small field from the cortex is shown. Capillary blood vessels in different stages of hyalinisation. At A, A, A, note persistent stenosed lumina. In the thickest vessel depicted the laminated appearance is well seen and necrotic nuclear material can be made out in the interstices. From a very elderly woman. About 300X.
TYPICAL HODGKIN'S DISEASE - LYMPH NODE

Field drawn from sub-capsular periphery. The full thickness of the capsule is not depicted.
Note - Fibrosis - Capsule, fibrils, fibrillae, and coralline deposit.

- Hodgkin giant-cells.
- Morbid mononuclear reticulum cells.
- Eosinophiles, plasma cells, neutrophiles.
- Rare individual cells are necrotic.
- Variation in size and staining of lymphocytes.

Serial 89. About 700X. H&E.
BASIC EXPRESSIONS OF FIBROSIS IN HODGKIN'S DISEASE

Fig. 1. Fibrillae. From a dissociated (macerated) specimen; fine collagenous filaments are seen. About 350X Van Gieson.

Fig. 2. Fibres. About 350X Van Gieson.

Fig. 3. Coralline deposits. (Compacted fibres; sometimes vascular hyalinisation also produces these formations). About 350X Van Gieson.

Fig. 4. Symplasmic transformation. Collagen fibrillae and small angular deposits form in the peripheral cytoplasm of epithelioid type reticulum cells. About 350X Van Gieson.
BASIC EXPRESSIONS OF FIBROSIS IN HODGKIN'S DISEASE

1.

2.

3.

4.
MORBID RETICULUM CELLS IN HODGKIN'S DISEASE

Fig. 1. Normal reticulum cell, nucleus and cytoplasm. (A small lymphocyte is drawn to scale for comparison).

Fig. 2. Normal reticulum cell; from a germinal centre, notching of the nuclear membrane is usual in this situation.

Figs. 3-9. Morbid reticulum cells which are still uninucleate. Increase of size of the nucleus is the most constant feature. Plasmasomes are fairly common, also evidences of necrosis.

Fig. 8. depicts two adnate epithelioid types, collagen formation by symplasmic transformation is indicated.

Figs. 10-12. Binucleate morbid reticulum cells, mirror-image form is common.

Figs. 13-24. Hodgkin giant-cells. These are all derived from a single section and shew the remarkable pleiomorphism of these cells.
MORBID RETICULUM CELLS IN HODGKIN'S DISEASE
MORBID RETICULUM CELLS IN HODGKIN'S DISEASE
MORBID RETICULUM CELLS IN HODGKIN'S DISEASE
FORMATION OF COLLAGEN FIBRES BY HODGKIN GIANT CELLS AND MORBID MONONUCLEAR RETICULUM CELLS

1. From Serial 101.
2. From Serial 122.
1. Direct metamorphosis in loose lymphatic tissue. The nuclei of the eosinophiles are single and correspond closely to large lymphocytic nuclei.
   About 750 X H&E

2. Lymphocytoid eosinophiles in the marginal sinus of a normal lymph node.
   About 500 X H&E
SELECTION OF LYMPHOCYTOID EOSINOPHILES

Note the amphophilic character of granules also the disparity of size and density. A small lymphocyte is shown, top left. Serial 79. H.P. H&E.
IMMATURE LYMPHOCYTES IN HODGKIN'S DISEASE

A survival islet of mature small lymphocytes is seen top left. It is separated from the new tissue by a fibril of collagen. The lymphocyte series cells in the tumour are larger and their nuclei are leptochromatic and tending to a vesicular type. (Lymphocyte cytoplasm was not discernible in this preparation).

Serial 110. About 600 X H&E.
FOLLICULOID DISPOSITION OF LATE HODGKIN'S DISEASE

Half-tone drawing. The black tissue is the cellular moiety, the pale grey the collagenous.  
Serial 156. Camera lucida X 5.
INCLUSION OF PHYSIOLOGICAL LYMPHOID TISSUE IN ESTABLISHED HODGKIN'S DISEASE

A late lesion is depicted; at A and B small unaffected lymph nodes are present.

Serial 199. Camera lucida X 5. H&E.
HODGKIN'S DISEASE, TYPICAL. (EARLY)

Cervical lymph node, at the periphery; the full thickness of the capsule is not drawn. All cells of the complex are represented, necrosis is limited to individual cells. Note the large (immature) lymphocyte nuclei.

Serial 89. About 600 X H&E.
HODGKIN'S DISEASE, CELLULAR TYPE

The general pleiomorphism is retained or enhanced. Morbid reticulum cells are relatively increased but generally remain somewhat small. Hodgkin giant-cells are retained but large forms are unusual. Eosinophiles persist, fibrosis is tenuous and generally uniform. Mitoses vary but are often increased in number.

Serial 193. About 600 X H&E.
ATYPICAL HODGKIN'S DISEASE, TUMOUR-LIKE TYPE

The drawing is representative of almost the entire node. Morbid mononuclear reticulum cells are numerous, they are more uniformly scattered than in the typical lesion, Hodgkin giant-cells are small. Eosinophiles are absent, the fibrosis is chiefly in fibrillary form and is uniform. The general uniformity of the pattern is tumour-like.

Serial 154. About 500 X H&E.
DIFFUSE FIBROBLASTIC HODGKIN'S DISEASE

The field is from the most cellular portions of the lesion. The majority of the cells are fibroblastic and their nuclei correspond to the range between morbid mononuclear reticulum cells and fibrocytes. The collagen in fibrils and fibrillae is forming by symplasmic transformation and is more densely oxyphile than the ground cytoplasm. The unstained spaces are shrinkage phenomena and are almost universal in this specimen. At the top an Hodgkin giant-cell is seen, these are rare in this example. Eosinophiles, small lymphocytes, (some of them immature) complete the picture.

Serial 168. About 600 X H&E.
HODGKIN'S DISEASE, PECULIAR VARIANT
(LINK TO RETICULUM CELL SARCOMA)

This growth is quite peculiar. All features of typical Hodgkin's disease can be made out but the overgrowth of large morbid mononuclear reticulum cells and Hodgkin giant-cells is extra-ordinarily florid. Note the retention of eosinophiles.

Serial 44. About 600 X H&E.
HODGKIN'S SARCOMA

Cellular portion of lesion, very numerous Hodgkin giant-cells including many bizarre forms, rare lymphocytes, and pyknotic neutrophiles, no eosinophiles, morbid mononuclear reticulum cells and fibroblasts; the whole set in a fibrous mesh.

Serial 202. About 500 X H&E.
The tumour is uniform. The type-cell nucleus has a morbid reticulum cell structure. The cytoplasm is sparse and feebly stained (visible only round a small proportion of nuclei).

1, 2. Small lymphocytes. 3. Necrotic lymphocyte nucleus.

4, 5. Mitoses.

6-9. Tumour reticulum cell nuclei. Note karyosomes and plasmasomes.

10, 11. Collagen fibrils.

Serial 109. About 600 X H&E.
Dichotomy of cell type characterises this tumour. Cells identical with the morbid reticulum cells of Hodgkin's disease are scattered uniformly through the growth. The majority of the cells are imperfectly differentiated to lymphocyte precursors. (Although not depicted here fibrosis and rare Hodgkin giant-cells were present).

1. Typical pathological reticulum cells.
2. Reticulum cells shewing affinities to lymphocyte series.
3. Lymphocyte precursors.

Note - Plasmasomes in some cells which are predominantly lymphocytic in character.

Serial 41. About 600 X H&E
LARGE CELL LYMPHOSARCOMA

Under low power the tumour appears an homogeneous cellular neoplasm in which the type cell is a young lymphocyte series cell, (lymphoblast). Under high power scrutiny this uniformity is lost, typical pathological reticulum cells are revealed, some of these cells are engaged in abortive fibre formations. Note how most of the lymphocyte precursors have karyosomes but some have plasmasomes. Several tumour cells are shewn in a capillary.

Serial 123. About 600 X H&E.
A DIAGRAM OF A DISSECTION OF THE LEFT AXILLA TO SHOW THE LOCATION AND DISPOSITION OF THE AXILLARY LYMPH NODES (TAYLOR AND NATHANSON NOMENCLATURE).

1. Axillary vein group.

2. Subscapular vein group.

3. Lateral thoracic vein group.

4. Thoracico-acromial vein group.

SPECIAL NECROPSY SERIES

LOCALISATION OF LESIONS

KEY TO THE DIAGRAMS

1. Normal lymph nodes, including those which are atrophic or shewing physiological reactive change.

2. Nodes absent, or at least not found.

3. Nodes shewing recognisable Hodgkin's disease (not necessarily typical).

4. Nodes shewing Hodgkin's sarcoma or reticulum cell sarcoma.

Note. Liver, spleen or other organs may be outlined in blue, purple, or red, indicating absence of lesion or participation in the morbid process. Important extra-nodal loci are similarly indicated.
LOCALISATION OF LESIONS
NECROPSY 'B'  AGNES MCNIVEN  P.M. 201:49 G.R.I.

LOCALISATION OF LESIONS
LOCALISATION OF LESIONS
LOCALISATION OF LESIONS
Necropsy 'H'  Mrs. Catherine Cameron  P.M. 155:52. G.R.I.  
Localization of Lesions
NECROPSY 'I'  MRS. MALA MOYES  P.M. 119:52. G.R.I.

LOCALISATION OF LESIONS
LOCALISATION OF LESIONS
LOCALISATION OF LESIONS

Necropsy 'K'  MARIE TRAYNOR  P.M. 415:53. G.R.I.
RESULTS OF TRYPSIN BLUE INJECTION IN RATS

(Series A., B., D. and E.) The death date of each animal is indicated by the position of the appropriate series symbol. A red ring round this indicates that tumour-like nodules were present in the liver.
TRYPAN BLUE.

DOSAGE

MgM

I.P.I.  25- 250- 200  15- 150- 100

RAT

A1  A2  A3  A4  A5  A6  A7  A8

TIME

DAYS

400

300

250

200

150

100

50

10
SERIES B.

I.P.I. DOSE TRYPAN BLUE.

TIME DAYS

250

300

200

150

100

50

0

RAT

B1 B2 B3 B4 B5 B6 B7 B8
S E R I E S  D.

TRYPAN BLUE.

I.P.I. DOSE MGM

TIME DAYS

DAYS

RAT

D1 D2 D3 D4 D5 D6 D7
RESULTS
WEIGHT RECORDS OF
SERIES A., B., C., D. and CONTROLS

RECORD OF AVERAGE BODY WEIGHTS.

WT GMs

0 10 20 30 40 50 60
TIME WEEKS

A
B
C
D
Co
Drawing of cut surface of rat liver showing tumour-like nodules and areas of necrosis.
Rat A.4 282 Days.

X4.