Review article

Some aspects of the metastatic process*

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The ability to metastasise can reasonably be regarded as the single most important characteristic of the vast majority of malignant tumours. It is also the least understood. Purely descriptive accounts of some of the most critical phases of the metastatic process are still lacking, and knowledge of the underlying mechanisms is rudimentary. Some advances are, however, being made and this review is concerned with work reported (for the most part) in the last decade which pertains to the two major metastatic pathways – lymphatic and haematogenous spread. The emphasis is on human studies but experimental data are included wherever necessary.

Spread of tumour in the lymphatic system

The lymphatics form the major route of dissemination for carcinomas and for many melanomas, neuroblastomas and malignant teratomas. Lymph-borne metastases from sarcomas are uncommon but are most likely to be encountered in alveolar and embryonal rhabdomyosarcomas, synoviosarcomas and epithelioid sarcomas.1

Penetration into the lymphatic system

Primary tumours do not contain lymphatic vessels and the main points of entry of malignant cells into the lymphatic system are lymph capillaries near the growing edge of the neoplasm. The microanatomical location and the density of lymphatics vary considerably in different tissues and are important determinants in the eventual incidence and distribution of regional lymph node metastases. Small lymph vessels resemble blood capillaries in their histological structure, consisting of endothelial cells linked by various types of interendothelial junctions and bounded externally by a basement membrane which is frequently discontinuous. Lymphatics from different parts of the body vary in the numbers and organisation of interendothelial junctions and the arrangements of the basement membrane2; regional differences of this kind may in part determine the ease with which the lymphatic system is invaded at certain anatomical sites.

The process whereby tumour cells penetrate local lymphatic vessels is ill-documented. The scanty information available is based on experimental studies with various types of tumour cells inoculated into the footpads of rats.3-4 Serial examination of tissues in the electron microscope indicates that the injected tumour cells first align themselves along the lymphatic channels. They then appear to protrude cytoplasmic processes between endothelial cells and migrate through interendothelial junctions, either as single cells or as small clumps. Damage to contiguous endothelium and perivascular collagen fibres is usually minimal or absent. Lymphatics in the rat footpad appear to be almost devoid of basement membrane, and the endothelium provides the sole (and manifestly ineffective) barrier to be traversed by invading tumour. The processes whereby neoplastic cells breach the endothelial layer are unknown, but possible factors include mechanical pressure exerted by motile tumour cells and the local release of substances which cause endothelial retraction or inflict minute lesions on or near the junctional complexes.

Once tumour cells, singly or in clumps, have crossed the lymphatic wall, they enter a low-pressure flow system in which circulation is maintained by a combination of intrinsic contractions of the lymphatic walls and extra-lymphatic pressure exerted by the movements of local muscles and transmitted arterial pulsations.5 Normal lymph contains little or no fibrinogen and platelets which, in the blood vascular system, play an important part in the arrest of circulating tumour cells and their subsequent escape into the extravascular compartment. Intralymphatic tumour is a common finding in surgical and necropsy material, usually seen as small clumps of cells lying free or occasionally still attached to the endothelial surface. Long segments of lymphatic vessels may be distended with growth which can sometimes be identified macroscopically. Intralymphatic tumour is

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difficult to recognise in poorly fixed, densely fibrous tissues where the lymphatic endothelium has been shed and the stroma has retracted to form small artefactual spaces round clumps of infiltrating tumour. Although important mainly as an indication of possible spread to regional lymph nodes, intralymphatic tumour without detectable nodal metastases is itself a notable finding for tumours at certain sites. It has, for example, been reported that patients with breast cancer with tumour cell emboli within intramammary lymphatics but without spread to axillary lymph nodes show an increased risk of developing visceral metastases.

**RETFENTION IN REGIONAL LYMPH NODES**

Disseminating tumour, carried in afferent lymph as single cells or small emboli, is initially trapped in the sinus system of the draining lymph nodes, the complex arrangement of which is strikingly revealed in low-power scanning electron micrographs. The retention of tumour cells in the sinus system has been confirmed in experimental studies in which neoplastic cells, sometimes labelled with radioisotope, are injected directly into afferent lymphatic vessels and their distribution followed for several days. There is, however, evidence from some investigations that tumour cells injected into local lymphatics may quickly traverse the regional lymph nodes. The efficiency (or otherwise) of the barrier function of lymph nodes remains unclear, and many aspects of the retention process are speculative.

**INTRANODAL TUMOUR: THE EARLY PHASES**

Various events ensue once tumour cells are retained in the local nodes – cell destruction, cell dormancy, and establishment and growth. The first two contingencies are scantily documented in experimental animals only, and the need for more detailed quantitative studies is obvious. The local conditions necessary for the establishment of a growing focus of tumour cells in a lymph node are obscure. Acquisition of an adequate blood supply is one critical requirement, and the normal lymph nodes have a well developed dual vasculature through hilar vessels and transcapsular anastomoses. The vasculature is adaptable and is rapidly augmented during, for example, the course of an immune response. Most nodal metastases appear to be adequately vascularised and may grow to form large masses several centimetres in diameter. Necrosis is, however, a characteristic feature of rapidly proliferating nodal deposits (from tumours such as melanomas, small cell anaplastic carcinomas of the bronchus, neuroblastomas, anaplastic seminomas and malignant teratomas) and of malignant lymphomas. Haemorrhage and necrosis may result from invasion of intranodal blood vessels by growing tumour. High endothelial-lined vessels are a characteristic (and diagnostically useful) feature of nodes infiltrated by T cell lymphomas, perhaps reflecting the production of T cell-associated angiogenesis factors. Growth of nodal metastases is usually continuous with progressive replacement of the pulp. The growth pattern and degree of cellular differentiation of the primary neoplasm tends to be reproduced, though the development of metastases which are less differentiated than the primary growth is a familiar finding. Undifferentiated nodal metastases may present diagnostic problems, particularly if the primary tumour is occult, and immunoperoxidase and other modern histological techniques are valuable. Large nodal deposits are often still confined within an intact capsule. Transcapsular spread is, however, a common feature for some tumours such as malignant melanomas and squamous carcinomas of the head and neck; capsular infiltration in partially involved lymph nodes is frequently seen, reinforcing scepticism for the simple notion that the capsule is disrupted by increased intranodal pressure.

**INTRANODAL TUMOUR: SOME LATE EFFECTS**

As the lymph node is replaced by metastatic tumour, lymph perfusion through the pulp will diminish and tumour cells will be carried to other nodes in the same or contiguous anatomical groups. Various complications of intranodal lymphatic obstruction will ensue. Retrograde lymph flow can cause tumour cells to be carried to more distant parts of the lymphoid system where they give rise to anatomically anomalous deposits of metastatic tumour. Retrograde flow is also responsible for spread of tumour cells, still within lymphatics, to sites outside the conventionally defined lymphoid system. Necropsy studies of disseminated breast cancer have shown widespread lymphangitis carcinomatosa in the visceral and parietal pleura and lung parenchyma. Similar lymphatic spread may occur in the liver. Local obstruction of lymphatic vessels will lead to oedema and, in the case of diaphragmatic lymphatics, to ascites – though the relative importance here of decreased lymphatic absorption and increased production of intraperitoneal fluid is uncertain. Tumour cells may disseminate from lymph nodes into the vascular system at several points. They include small intranodal vessels; extranodal vessels invaded by metastatic tumour which has breached the nodal capsule; lymphaticovenous communications opening up in the involved lymph node; and the thoracic duct. Such spread is usually regarded as a late event, occurring when most or all of a lymph node is replaced by metastatic tumour. The basis for this assumption is unclear, and there are no a priori reasons why
intranodal tumour cells should not disseminate further in the blood stream at an earlier phase.

The localisation of nodal metastases is largely determined by the local anatomy, though anomalies sometimes arise. Normal pathways of lymphatic drainage may be modified by previous radiotherapy and major surgery. Retrograde spread of intranodal extracranial metastases has already been noted. Certain aspects of the basic anatomy at some sites are still unclear, and the continuing discussion as to whether each breast quadrant does or does not have a consistent and specific pattern of lymph drainage provides one example. The more important question of predicting whether nodal metastases will develop – as opposed to forecasting their anatomical distribution – is complex and speculative, and falls outside the scope of this short account.

**Spread of tumour in the vascular system**

Sarcomas disseminate predominantly by the blood stream, and melanomas, neuroblastomas and malignant teratomas spread almost equally in the blood and lymphatic systems. Haem dissemination is an important and often underestimated metastatic route for carcinomas, even though lymphatic spread predominates in many instances.

**Penetration into the blood vascular system**

Malignant cells enter the circulation by invading blood vessels situated either within the substance of the main tumour mass or near its advancing edge; they may also gain access indirectly through the lymphatic system – see earlier. The vasculature of tumours originates from pre-existing vessels in the tissue of origin and from newly formed vessels derived from them. The stimuli for angioneogenesis within tumours are unclear, but they are likely to stem from host sources (especially local lymphocytes) as well as from neoplastic cells themselves. Such vessels are extensive and often imperfectly formed with irregular lumina, discontinuous and sometimes abnormal endothelium, defective basement membrane and (in larger vessels) scanty or absent perivascular connective tissues. Sarcomas often contain large blood-filled channels wholly or partly lined by malignant cells which can shed direct into the blood. Vascular endothelium in experimental tumours is capable of intense proliferative activity and, in clinical material, endothelial hyperplasia is a distinctive feature of high grade astrocytomas and (to a lesser extent) other primary intracranial neoplasms. Similar vessels are seen in some brain metastases, and they have also been described in the very rare extracranial metastases from gliomas in sites such as cervical lymph nodes. Despite the extensive vasculature, the rapid endothelial turnover and the overall high perfusion rate in many tumours, regional perfusion with them is frequently defective. Many of the abnormal vessels are excessively permeable and the interstitial compartment in most tumours is large. The accumulation of pericellular fluid is enhanced by the absence of any intrinsic lymphatic drainage. Regions of necrosis within a tumour, and the accumulation of oedema fluid and fibrin at its edges, are at least in part explained by such circumstances. Necrosis is often marked in rapidly proliferating tumours with a large cell loss fraction, and such tumours, which are sometimes very small, have a notably high metastatic rate. Examples include small cell anaplastic carcinomas of the bronchus, melanomas, malignant teratomas and choriocarcinomas. There is some experimental evidence that local necrosis favours the detachment of cells from the growing tumour mass.

Normal blood vessels at or near the edge of the tumour are fewer and less readily penetrated by invading cells, but they represent the major route for haem dissemination. Capillaries and small veins are the principal sites of entry. The process whereby tumour cells cross the vessel walls is unknown. Larger vessels, occluded by neoplastic cells and thrombi, are usually easy to recognise but smaller vessels lacking organised connective tissue coats are more problematic, particularly in dense stroma. Immunohistochemical stains for Factor VIII-associated antigen in normal endothelium are helpful in suitably fixed tissues. Invasion of large, well-perfused, thick-walled veins with contractile muscle coats appears to be particularly hazardous. An example is provided by recent work on venous invasion by carcinoma of the rectum in which close correlation was demonstrated between infiltration of extramural, thick-walled veins and the incidence of hepatic metastases. Local arteries are rarely invaded by neoplastic cells. Infiltrating tumour characteristically encircles them but appears to stop a short distance beyond the outer border of the adventitia. The thickness or composition of the arterial wall cannot be implicated, and there is some experimental evidence which suggests that transmitted pulse pressure may be responsible.

**Circulating tumour cells**

Tumour cells free in the circulation have been extensively studied, but to little ultimate benefit. Most of the work is rendered suspect by inadequate attention to technical details (number and size of blood samples, their timing, sites of venepuncture, choice and methods of cell concentration) and to the criteria for identification of malignant cells in
cytological preparations. Such investigations are – inevitably – wholly qualitative and certain simple questions are left unanswered: What proportion of circulating cells is alive? How many cells are still able to divide? How many cells lack the vital capacity to stick to vascular endothelium? The demonstration of neoplastic cells in leucocyte concentrates appears to have little or no prognostic implication for any tumour at any site so far studied.

Some quantitative data on the release of cells into the circulation from transplanted tumours are available from studies in mice and rats, and extensive work has been done on the fate of tumour cells injected intravenously in experimental animals. Such manoeuvres are convenient but artificial, in that they by-pass the essential phase of vascular invasion, and extrapolation of results to the metastatic process as it really exists is often questionable.

**ARREST OF CIRCULATING TUMOUR CELLS**

The morphological changes which ensue when tumour cells are arrested in the circulation were meticulously described in both clinical and experimental material in the early part of this century. In brief, tumour cell emboli in small vessels of the lungs, liver and other tissues stick to vascular endothelium where they are covered by fresh thrombus. Some tumour cells grow progressively, filling the lumen and eventually escaping out into the extravascular compartment. Others remain entrapped in thrombi which are endothelialised and become organised with subsequent hyalination and fibrosis. Organising thrombi occasionally contain a few well-preserved tumour cells, some of them even seen in mitosis. These morphological events, based on sequential but inevitably static observations, were confirmed by Sumner Wood in his classical studies of tumour cells injected proximal to ear chambers implanted in rabbits. The process is one of great complexity, and several aspects remain controversial. Examples include the interplay of mechanical, haemodynamic and electrostatic forces in the initial impact of tumour cells on the endothelial surface; the processes whereby contact becomes attachment; the initiation of local thrombus formation and (by implication) the role of anticoagulants and fibrinolysins; and the processes whereby tumour cells extravasate into the surrounding tissues. These topics are discussed elsewhere but one general point needs to be stressed. Morphological studies in the light microscope, ultrastructural investigations and the direct cinemicrographic observations made in the living animal all emphasise that tumour cells sticking to vascular endothelium do not invariably complete the full sequence of vascular destruction and extra-vacation. Many tumour cells will die within the thrombus, and others will be swept back into the circulation when the surrounding platelet-fibrin aggregates dissolve. No quantitative data are available, but it is reasonable to suppose that only a minority of arrested tumour cells will succeed in escaping into the interstitial compartment – metastasis, for all its lethal consequences, is in biological terms a distinctly inefficient process.

**Use of in vitro models**

Some of the mechanisms whereby tumour cells escape from blood vessels have recently been investigated with in vitro systems. Bovine aortic endothelium, which also synthesises basement membrane components, provides a useful substrate; and there is growing knowledge of the biochemistry of basement membrane components, notably the adhesive glycoproteins (such as laminin and fibronectin) and type IV collagen. A summary of some recent work is given in Table 1. The results are interesting but certain limitations are immediately apparent. The tumour cells so far tested are somewhat limited in variety and are derived from cell lines. The *dynamic* interactions between tumour cells and vascular endothelium are virtually eliminated. Three-dimensional vascular structures are reduced to a simplified two-dimensional planar surface or a pure chemical substrate. Other components, notably local platelet-fibrin aggregates, are absent. Recent experiments with artificial blood vessels built up from bovine aortic endothelium and multilayers of rat smooth muscle provide some illuminating results. Destruction of subendothelial smooth muscle by human fibrosarcoma cell lines appears to be retarded in this system if intact endothelium is present – a finding which would be quite inapparent in a simpler (but arguably less representative) test system. Considerable caution is clearly necessary in extrapolating from static in vitro models to the turbulent situation that obtains in vivo.

**EXTRAVASCULAR TUMOUR**

Early events after tumour cells have crossed into the extravascular space are particularly ill-understood. Many cells are likely to die for purely metabolic reasons while others may be destroyed by host cells which initially will be at a numerical advantage. Some extravascular tumour cells will probably enter local lymphatics. Establishment and growth beyond a critical surface area/volume relationship is likely to be determined by the acquisition of a fibrovascular stroma. Wood’s cinemicrographic studies showed that a developing nidus of extravascular tumour in the rabbit ear was well vascularised within 36 h. Micrometastases lacking an adequate blood supply are
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Table 1  Aspects of extravasation of tumour cells studied in in vitro systems

<table>
<thead>
<tr>
<th>(a) Penetration of endothelium</th>
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<tbody>
<tr>
<td>1 Loose attachment of tumour cells to endothelium</td>
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<tr>
<td>2 Retraction of endothelium with exposure of basement membrane</td>
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<tr>
<td>3 Spread of tumour cells beneath endothelial monolayers.</td>
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<tr>
<td>- Complete process 1–3 demonstrated with tumour cells derived from cell lines from human, rat and mouse tumours.</td>
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<tr>
<td>- Incomplete process 1–2 demonstrated with various normal control cells EXCEPT for human polymorphs and monocytes.</td>
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<tr>
<td>- Capacity to attach to endothelial monolayers in vitro varies with different tumour types, possibly related to metastatic potential</td>
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<th>(b) Adhesion to basement membrane</th>
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<tr>
<td>Tight attachment of tumour cells to adhesive glycoproteins such as fibronectin, laminin and (?) type IV collagen (codistributed with laminin).</td>
</tr>
<tr>
<td>- Demonstrable by immunofluoresence and immunoperoxidase methods</td>
</tr>
<tr>
<td>- Capacity to attach to basement membrane-derived glycoproteins in vitro varies with different tumours</td>
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<tr>
<td>(? carcinoma cells binding to laminin, sarcoma cells to laminin and fibronectin).</td>
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<tr>
<th>(c) Destruction of basement membrane</th>
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<tr>
<td>Measured by degradation of whole basement membrane or purified basement membrane-derived type IV collagen.</td>
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<tr>
<td>Destructive activity is:</td>
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<tr>
<td>- substrate specific</td>
</tr>
<tr>
<td>- associated with tumour cells and their supernatant culture media</td>
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<tr>
<td>- probably related to metastatic potential</td>
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Data based on references 36–45.

likely to remain dormant; but the whole question of tumour dormancy remains obscure and the relative contributions of neovascularisation, tumour cell kinetics and host inflammatory and immune responses are almost totally speculative.

Once a nidius of tumour cells starts to expand it will infiltrate surrounding tissues in the same way as a primary neoplasm. The topic of tumour invasion falls outside the scope of this review, but one point should be noted – the growing evidence that host elements play an important role in the invasive process. In soft tissues, for example, it is local polymorphs, macrophages and fibroblasts near the tumour’s growing edge which elaborate potent proteases. Comparable activity in the tumour cells themselves may be less or even absent though the operation of functional interactions between neoplastic cells and the tumour-associated stromal components is virtually certain.

A comparable situation is seen when tumour cells are invading cortical bone: most bone destruction appears to be mediated by osteoclasts which are activated locally by both the tumour and the adjacent stroma.

Human tumours implanted as xenografts in immune-depressed animals are unsatisfactory models for studying local growth patterns. Irrespective of their previous clinical behaviour, such tumours form slowly growing lesions enclosed in fibrous tissue. Neoplastic infiltration of local tissues, and metastasis, are both exceptional events. The later stages of metastatic development need no detailed comment. Growth is usually progressive and, in some instances, doubling times have been shown to be greater than those of the primary tumour by a factor of 1.5 to 2. Spontaneous arrest and regression of metastases are exceptionally rare and the underlying mechanisms are unknown. Most established metastases resemble their parent tumour in histological structure though, as in lymph nodes, marked dedifferentiation may lead to diagnostic problems for the pathologist. Identification of small lesions, especially in the liver and bone marrow, is aided by immunoperoxidase techniques and the widely distributed epitheplium membrane antigen (EMA) is proving particularly useful.

Localisation of metastases

The distribution of lymph-borne metastases was discussed earlier and this final section is concerned with the localisation of blood-borne tumour. Some of the factors implicated are summarised in Table 2.

The large pulmonary and hepatic vascular beds form the first and quantitatively most important filters for trapping circulating tumour cells, and the lungs and liver are the two commonest sites of haemic metastases. A third set of primary filters is probably provided by the terminations of Batson’s para-vertebral venous plexuses in parts of the axial skeleton. The importance of these plexuses as a metastatic pathway is disputed and the situation is complicated as blood-borne tumour cells may reach the skeleton by at least two routes – directly through the paravertebral venous plexuses and also by the systemic circulation through the peripherally directed branches of the nutrient arteries. Tumours also invade contiguous bone by direct extension. Bross and his colleagues have proposed that the lungs and liver act as “key generalising sites” from which subsequent metastases are generated. Their data are
now extensive and are derived from large numbers of tumours arising at many sites, but the limitations of the hypothesis need to be borne in mind. The information is based on necropsy findings and thus reflects advanced, often long-standing, end-stage disease which has been previously treated. Lymphatic dissemination is not included and it was noted earlier that parenchymal organs such as the lungs and liver can be extensively infiltrated by lymph-borne tumour from a distant primary in the form of lymphangitis carcinomatosa. It is self-evident that a proportion of tumour cells initially trapped in the primary capillary filters can escape. In the early stages, some cells may squeeze through capillary lumina and enter the systemic circulation while others traverse vascular shunts. Later, tumour cells may be released from established foci in these regions; Willis'17 has stressed that such deposits may be minute.

The non-random distribution of systemic metastases outside the lungs, liver and parts of the axial skeleton is one of the oldest observations of tumour pathology. It still remains true, but certain features may be noted. Detailed necropsies on some patients with tumours such as carcinoma of the breast, melanoma and some soft tissue sarcomas, show that metastases can develop at almost any site. Metastases in the spleen are not the rarities they are sometimes made out to be.17 Secondary deposits in the adrenals are probably drawn from a wider spectrum of primary tumours than is often supposed.60 Furthermore, metastatic patterns from any one tumour type are not immutable. Necropsy studies are now showing that squamous carcinomas of the head and neck quite frequently metastasise to distant sites,61 that osteosarcomas may also spread to extrapulmonary tissues such as lymph nodes, abdominal viscera and other parts of the skeleton62 and that the distribution of metastases from breast cancer may be changing as a result of treatment.63 Some of the factors that are implicated in the localisation of blood-borne metastases have been summarised in Table 2. The traditionally disparate elements of the "haemodynamic" and "soil and seed" theories of metastatic localisation, associated with James Ewing and Stephen Paget, have been deliberately merged as the distinction seems futile.

Apart from the lungs and liver, no wholly consistent association has yet been established between the organ distribution of systemic blood-borne metastases and regional haemodynamic characteristics such as blood volume, blood flow and transit times within a specific organ.64 The methods available may not be sufficiently accurate and the importance of more refined haemodynamic studies, particularly with respect to regional variations in blood flow within a viscus, remains to be examined. Local variations in vascular structure and function have received little attention. Earlier workers65 suggested that thick-walled arterioles in tissues such as the spleen and skeletal muscle might provide a mechanical barrier preventing the egress of circulating tumour cells, but other more subtle features may operate as well. The interaction of local haemodynamic factors and local vascular organisation is well illustrated in the bone marrow. The sinusoidal system in red marrow has a large blood volume and slow perfusion with a tendency to intermittent reflux flow, and the spaces are lined by a discontinuous, often attenuated endothelial layer which appears to lack tight junctions.66-68 Such conditions are well suited to ensure (under physiological conditions) the unimpeded passage of mature blood cells from extravascular haematopoietic foci in the marrow into sinusoidal blood. These same conditions will, however, be no less favourable for the reverse passage of tumour cells from the sinus capillaries and their subsequent colonisation of the extravascular tissue spaces. The normal tissue milieu may be modified equally by local and general systemic factors affecting the whole body such as inflammation, trauma, irradiation and immune status. There is experimental evidence, for example, that immune status will modify the retention, distribution and growth of disseminated malignant cells.69-71 The complexity of the situation is illustrated by recent experience with athymic nude mice which are almost refractory to developing lung deposits from allogeneic or xenogeneic tumour cells even when injected directly into the venous system.72

The extent to which the distribution of metastases is determined by tumour cells themselves is unknown. Factors such as the rate of release of cells from the primary lesions, and their presence in the circulation as single cells or clumps, are likely to be important — but the fundamental question relates to whether certain intrinsic properties of the tumour cells determine their tendency to seed "preferentially" in certain sites or, indeed, to metastasise at all. The evidence available is based on experimental studies, and the implications for man are unclear. A detailed appraisal of this work falls outside the scope of the present review, but two of the principal strands of evidence may be very briefly summarised as follows:

1 A series of transplantable mammary carcinomas in rats has been established by Kim73 with consistent and distinctive metastatic patterns. Blood-borne and lymphborne metastases may develop and the target sites include the skeleton in two instances. Investigations suggest that the metastatic variants have unstable plasma membrane structures with abnormal shedding of glycoproteins, glycolipid
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antigens, surface marker enzymes, glycosyltransferases and other enzymes. Such abnormalities may lead to alterations in immunogenicity, membrane permeability and to cell locomotion; but the extent to which such features may determine the differential localisation of metastatic tumour cells in target organs remains unclear.

2. Extensive studies have been made by Fidler and his colleagues on the metastatic potential and organ selectivity of transplantable tumours in mice, notably the B16 melanoma. Sublines of the tumours have been established with an enhanced capacity to grow not only in the lungs — the usual site of metastasis for the B16 melanoma — but also in the brain, ovary and liver, following intravascular injection of cells at an appropriate site. The isolation and cloning of high- and low-metastasising strains of cells provides unequivocal evidence in favour of specific metastatic subpopulations. There are difficulties in the appraisal of experiments in which tumour cells are injected directly into the vascular system, and it is essential to confirm the behaviour of these seemingly selective tumour cell lines by growing them as subcutaneous or intramuscular implants from which "natural" metastases may be generated. Evidence from such experiments is conflicting, but the distinction between colonising and truly metastasising tumour cells which seems to be emerging illustrates the interpretative difficulties of this kind of modal system.

There is growing evidence that human tumours are heterogeneous with respect to features such as karyotypes and DNA content, the presence of hormone receptors, antigenic determinants and cell surface constituents, pigment synthesis, drug sensitivity, in vitro growth characteristics and growth as xenografts in immune-suppressed mice. Similar differences, albeit less clearly defined, have been described for some properties from primary and metastatic tumours in man. Whether such characteristics can be associated with specific genotypic sub-populations with an enhanced capacity to metastasise is wholly unknown, and the need to consider the probable operation of several different, non-exclusive metastatic mechanisms should require no special pleading.

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