Recent advances in the study of tumour invasion and metastasis

D Tarin, Y Matsumura

Introduction
The application of recently developed research techniques to the study of tumour invasion and metastasis has uncovered new and surprising information which has focused increasing clinical and scientific interest on this aspect of neoplasia. The magnitude of the effect of this problem on health is revealed by reference to the annual mortality statistics published by the Registrar General of England and Wales. These show that about one in three of the population die of the consequences of metastatic cancer, or are found to harbour asymptomatic metastatic tumour deposits at necropsy. Research on this subject is therefore directed at controlling a major, clinically important phenomenon. A selection of recent advances which offer new opportunities for early diagnosis or treatment of this aspect of malignant disease are surveyed in this brief overview.

Clinical and experimental background
The invasive and metastatic activity of malignant tumours has, until now, seemed to be an almost insuperable problem, best illustrated by a clinical example. The patient whose leg is shown in fig 1A had initially presented two years earlier with a small melanoma. This was treated by wide local excision and grafting of skin from the back of her torso. Subsequently an advancing wave of dermal metastases (fig 1B) gradually ascended her leg. Metastatic tumour then colonised and replaced her inguinal lymph nodes and went on progressively to take over the para-aortic group. Finally, haematogenous metastasis began, evidenced by the presence of pigmented tumour deposits in the brain at necropsy (fig 1C). Death, in this case, was due to an unrelated condition, but is more often due in disseminated malignancy, to organ failure following replacement by tumour (fig 1D). As the metastases in this patient only appeared after excision of the small primary tumour, they must have already been disseminated. Equally, the eventual appearance of metastatic deposits in the grafted skin (fig 1A), donated from a remote site which never developed dermal secondaries, illustrates that metastases themselves act as new foci for further dissemination, resulting in a geometric increase in the impact of the tumour on the host.

Clinical experience, therefore, teaches that malignancy is often already systemic from very early in the life history of a neoplasm and that patients may harbour many covert seeding tumours at the time of first presentation with symptomatic lumps. Diagnostic and therapeutic policy need to be predicated on this understanding. As the treatment of established metastases is presently rather crude, toxic, and limited in success, the clinical man-

Figure 1A Pigmented metastases in the leg of a patient with malignant melanoma. The site of resection of the primary tumour can be seen and deposits are even present in the skin graft covering the deficit. 1B Thigh of the same patient showing numerous cutaneous metastases, some of which are pigmented. A pale scar can be seen in the groin at the site of previous resection of deposits in the inguinal nodes. 1C Portions of two slices of the brain of this patient. Pigmented metastases can be seen in the grey matter of the cerebral cortex. 1D Unpigmented metastases in liver of another patient with malignant melanoma.
agement of malignancy will benefit from efforts to improve early diagnosis of cancer and to impede tumour dissemination. New advances which contribute in some way to these objectives may be grouped as follows:

1. The earlier a tumour can be detected, preferably non-invasively, the better because surgery and radiotherapy are now very effective for the treatment of neoplasms that are still localised. New strategies described below, of searching for tumour related abnormal gene activity with recently developed amplification methods to increase sensitivity, are therefore being developed to facilitate early detection.

2. Other work in progress offers the prospect of methods for the evaluation of the metastatic potential of a tumour biopsy or needle aspirate, to assist choice of treatment.

3. It has recently been demonstrated that tumour cells entering the circulation in patients with cancer do not necessarily form metastases even when they do so in large numbers. This has implications for judging prognosis and treatment and for monitoring disease recurrence.

4. Another group of results pertain to the aim of blocking tumour invasion and metastasis in patients who could already have some deposits. Such endeavour is worthwhile even when some metastases are known to be present, in the service of trying to gain the patient some advantage from reduction in the geometric advance of the disease.

Current understanding of the spread of cancer in the human body is based on a good deal of previous research which has established that firstly it is a stepwise phenomenon (fig 2) capable of being undertaken by only a subpopulation of cells within the heterogeneous masses of cells in a tumour, secondly that it is highly selective and results in massive losses among the cells which set out on the venture and thirdly that it is stably propagated over many generations among the cells which survive to form secondary tumour deposits. Hence it is likely to be genetically programmed. It is also known from many clinicopathological studies that metastasis can occur via several anatomical pathways and that the distribution of tumour deposits tends to occur in non-random, organ selective patterns according to the site and histogenetic origin of the primary tumour. It seems, therefore, to be subject to modulation by local environmental factors in the sites of tumour cell arrest, an interpretation also favoured by recent experimental data. The advances to be discussed are consistent with, and support, this earlier framework of understanding.

Early tumour diagnosis as a practical means of controlling metastatic sequelae

Recent studies have provided evidence that the activity of the CD44 gene is severely deranged in many types of tumours. This gene normally codes for a family of heavily glycosylated cell surface proteins, the multiple isoforms of which exercise many important cellular functions. In cancer there is chaotic overproduction of many unusual mRNA transcripts, relative to the picture seen in corresponding normal or non-neoplastic tissues. As the disorder is present in early tumours (Matsumura et al, unpublished observations) and can be identified in very small samples using amplification techniques, it seems a promising candidate to study as a possible marker for early diagnosis and for monitoring patients for local and distant recurrent disease.

The first observation that there is gross overexpression of numerous splice variants from this gene in human tumour tissue was serendipitously made in the course of studying metastatic cells to test the hypothesis that such cells are inappropriately activating genetic programmes for white cell (especially lymphocyte) recirculatory traffic. An earlier study in

Figure 2. The multistep process of metastasis.

The Multistep Process of Metastasis

1. Primary malignant tumour
2. Multiplication, invasion and vascularisation
3. Invasion of vessels
4. Entry of tumour cells into the cardiovascular system
5. Extravasation of tumour cells in a downstream organ
6. Multiplication and vascularisation of secondary deposits in distant sites
7. Arrest of tumour cells in downstream organs
8. Survival of high velocity collisions, and elusion of the immune system
this laboratory had provided evidence that the integrin VLA-4, commonly seen on trafficking lymphocytes, is upregulated on metastatic but not on non-metastatic tumour cells. It was therefore decided to examine CD44 expression in such cell lines, because it had been identified as a lymphocyte homing molecule, aiding these white cells to traffic through lymph nodes. The results with human cell lines were disappointing, there being no clear correlation with metastatic capability. However, a report from another laboratory, of differential expression of a particular small portion of the gene in metastatic and non-metastatic cell lines derived from a rat pancreatic adenocarcinoma indicated that it may play a part in this process in some circumstances. We therefore examined CD44 gene activity in fresh human tumour tissue and metastases with more sensitive techniques. The resulting findings were dramatic (fig 3). They were obtained with the technique of reverse transcription, followed by polymerase chain reaction amplification (RT-PCR), and showed very disorderly expression of this gene in tumour samples. It can be seen from this picture that the differences between tumour tissue and normal tissue were obvious and increased in severity with progression of malignancy to metastatic behaviour. Our subsequent studies have shown that such abnormalities are present in a wide variety of common cancers (table), but not in corresponding normal tissues. By refining the technique we have also been able to show that with this method even small numbers of tumour cells can be detected in urine obtained non-invasively, by natural micturition, from a very high proportion of patients with bladder cancer. The implications for non-invasive investigation of patients for the possible presence of tumours in accessible sites are obvious and are being explored. So far the results are scientifically very encouraging and interesting, but the numbers of cases with malignant disease and of controls with non-neoplastic conditions need to be expanded greatly, with inclusion of double blind protocols, before it will be possible to decide whether the method can be dependably used in clinical practice. Also, the work to date has shown that disturbed activity of the CD44 gene is not exclusively associated with metastasis, but is seen in many types of tumours and seems to increase progressively with malignancy. The exact reasons for such abnormal activity of this gene and of its association with neoplasia await further investigation.

Other laboratories have explored the possibility of detecting p53 or ras mutations with PCR based techniques, to diagnose the presence of cancer cells in the urine of patients with bladder cancer, but the methods needed for such DNA Based tests are more complicated and their usefulness for general cancer detection is limited by the fact that only a proportion of patients with the condition have the mutation in their tumour cells. Although an improvement in the methods of extracting RNA from clinical samples is greatly needed, RNA based diagnosis promises to be simpler and more practical for the investigation of a given individual who may currently have disease. DNA based techniques are probably more suited to the analysis of familial and aetiological associations. A cautionary note needs to be made: mRNA is notoriously susceptible to degradation by any

Table  Derangement of CD44 expression in tumours

<table>
<thead>
<tr>
<th>Type of tissue</th>
<th>Number of patients/ volunteers</th>
<th>Number showing increased splice variants</th>
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<tr>
<td>Neoplastic</td>
<td></td>
<td></td>
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<tr>
<td>Breast cancer</td>
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<td>46</td>
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<tr>
<td>Colon cancer</td>
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<td>21</td>
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<tr>
<td>Bladder cancer</td>
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<tr>
<td>Stomach cancer</td>
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<td>6</td>
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<td>Thyroid cancer</td>
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<td>1</td>
</tr>
<tr>
<td>Fibroadenoma</td>
<td>1</td>
<td>1</td>
</tr>
<tr>
<td>Prostate cancer</td>
<td>3</td>
<td>2</td>
</tr>
<tr>
<td>Non-neoplastic</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Normal breast</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>Cystic disease of breast</td>
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<td>0</td>
</tr>
<tr>
<td>Normal colon</td>
<td>9</td>
<td>0</td>
</tr>
<tr>
<td>Crohn’s disease</td>
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</tr>
<tr>
<td>Ulcerative colitis</td>
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<td>0</td>
</tr>
<tr>
<td>Appendicitis</td>
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<td>0</td>
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<tr>
<td>Normal bladder</td>
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<td>4</td>
</tr>
<tr>
<td>Peripheral blood leukocytes</td>
<td>10</td>
<td>0</td>
</tr>
<tr>
<td>Bone marrow</td>
<td>3</td>
<td>0</td>
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</table>
Looking for metastases

388

Figure 4

Experimental protocol for transfection of metastasis.

Transfection of Metastasis

1. Extract DNA
2. A375M DNA
3. Mixed DNA
4. DNA transfection procedure
5. TR4 cells: non-metastatic mouse cell line
6. Inject cells containing tagged DNA
7. Look for traces of human DNA by ‘Southern blotting’
8. Isolate and test fragments of human DNA for effects on metastasis by repeating steps 3 to 6

New methods for assessment of metastatic potential and prognosis

In a recent series of experiments we succeeded in transferring metastatic capability by transfecting total genomic DNA from a human malignant melanoma cell line into a non-metastatic mouse tumour cell line. The plan of experiment is summarised in fig 4. Secondary transfer of this property was also achieved in a second round of DNA transfection into fresh cells of the same non-metastatic mouse line. The DNA used for this second transfer was obtained from a cell line derived from a pulmonary metastasis made by the primary transfectants. Concomitant transfer of human DNA through both transfection cycles has been shown by several methods, including probing Southern blots with probes for human specific Alu repeat sequences, in situ hybridisation, and Alu PCR. Finally, a portion of the transfected human DNA was recovered by screening a genomic library made from the transfected metastatic cells. This has been sequenced and is a 2858 base pair fragment containing previously unrecorded human coding regions. Using PCR primers we designed to one of these, in an RT-PCR analysis of mRNA extracted from fresh human tumours and normal tissues, we have seen marked differences in the expression of this sequence in malignant tumours, relative to benign tumours and normal tissues. More samples are being examined but the results to date clearly indicate that the probe distinguishes tumour tissue with metastatic potential and that it could therefore be useful in clinical practice.

In 1988 Steeg and colleagues cloned and sequenced a separate gene which seems to be relevant to the metastatic process and may have diagnostic potential. This gene, designated nm23, was isolated by differential hybridisation from a non-metastatic mouse cell line and is conserved in human cells. Studies to date show a promising correlation between reduced expression of the gene and metastasis in patients with breast cancer, but the results in colon carcinoma have been contradictory. The gene might prove to be a useful marker and further results are awaited with interest.

Very recently results from three laboratories have indicated that counting the number of capillaries in the most vascularised area of a tumour might give a useful pragmatic indication of patient survival or of tumour recurrence in breast cancer and non-small cell lung cancer. The underlying basis for this clinical evaluation is the well established knowledge that vascularisation is essential for tumours to grow beyond a few millimetres in size, and for dissemination of tumours by blood and lymphatic routes. Although there is very good statistical correlation between vessel density in histological sections and prognosis, it remains to be seen in further studies whether it is a useful index for the evaluation of individual patients.

Invasion

Once tumour cells have activated internal genetic mechanisms for dissemination from the expanding primary lesion, they radiate outwards into the surrounding tissue releasing proteases, especially collagenases, (fig 5) and destroying much of the extracellular
matrix which they encounter\textsuperscript{35-37} (fig 6) until they collide with local capillary blood vessels and lymphatics. The former are sheathed in basement membranes which are not readily breached by most normal cells. However, they have to be traversed by tumour cells programmed to disseminate by entering vascular channels, or by patrolling lymphocytes passing through the tissues on routine surveillance missions.

These membranes contain laminin and collagen type IV and the latter is selectively digested by a specific metalloprotease, gelatinase, otherwise known as collagenase IV, which is reported to be secreted in increased quantities by several types of tumour cells. It has also been suggested that this enzyme is secreted in greater quantity by cell lines which are highly metastatic than by ones which are not.\textsuperscript{38} There is little doubt now that both this and collagenase I, which selectively digests collagen I, the main structural component of the extracellular mesenchyme, are active in tumour invasion and metastasis. Even so, the production of these lytic agents in tumours is spatially and temporally erratic, and there is no direct evidence that collagenase values of individual lesions are of diagnostic or prognostic value in clinical oncological practice. With the recent advent of new, collagenase specific inhibiting drugs, which remain active after absorption via the oral route, however, the possibility of reducing local tissue destruction or delaying distant metastasis has appeared on the horizon. Long term trials will be needed to evaluate such agents, not only to assess whether they are effective, but also because of the need to ensure that they have no detrimental effects on collagen metabolism and turn-over in the normal tissues and organs of the host. Potentially, such agents could be effective against local invasion and metastasis from the primary tumour as well as in impeding similar activities of secondary deposits.

Figure 5 Polyacrylamide gel electrophoresis of digestion products (arrows) of type 1 collagen exposed to culture supernatant fluids, from human breast carcinomas (tracks 2 and 4 numbering from left). Digestion was inhibited by addition of EDTA (tracks 3 and 5) confirming it was due to a specific mammalian metalloprotease—namely, collagenase. Track 1 undigested collagen control; track 6 molecular weight markers.

Tumour cell dissemination and patterns of metastasis

Over a century ago Stephen Paget realised that the distribution of metastases from such tumours is not random. He inferred that the process depended both on intrinsic properties of the disseminating tumour cells and of the organ in which they became detached after being scattered. This work gave rise to the "seed and soil" hypothesis of metastasis.\textsuperscript{39} Actually this deduction was a formidable achievement, based on remarkably clear, direct, and independent thinking, especially if one remembers that the bulk of the pathological establishment at that time, including Virchow, were in dispute about whether tumour metastasis resulted from dissemination of cancer cells or of infective agents released by the tumour.

Although this controversy was soon settled, direct confirmation of the hypothesis had to await experiments with cultured tumour cell lines in inbred syngeneic strains of animals and in nude mice. Final confirmation in humans took still longer and was only obtained through a fortuitous opportunity provided by the introduction of a novel form of palliative treatment for patients with intractable ascites caused by inoperable abdominal malignancy.\textsuperscript{12} The technique of peritoneo-venous shunting is sometimes used to relieve pain and discomfort in patients with recurrent malignant ascites. These patients go into severe metabolic imbalance if large volumes of such fluid, and its constituent nutrients and metabolites, are regularly removed by the simpler procedure of paracentesis. The shunt resolves the metabolic problem by

Figure 6 Electron micrograph of the advancing edge of a deeply invasive skin carcinoma showing extensive disorganisation of connective tissue extending several microns in advance of the neoplastic epithelium (E) which is extending clear pseudopodia into the vacant space created by lys of adjacent tissue. There are few remaining collagen fibres (arrows) and the basement membrane has been demolished.
returning the fluid to the circulation via the jugular vein, but simultaneously infuses billions of viable tumour cells, floating in the ascites, directly into the blood for protracted periods of time. Amazingly, some patients with metastatic cells were undoubtedly viable and tumourigenic on implantation in the peritoneum did not develop any detectable metastases in any organ, despite surviving for up to 2 years before dying from local tumour burden in the original site. Other patients developed seedling tumours via the shunt in some organs but not in others, and the organ distribution of these corresponded to that of known metastases before the shunt was inserted.

The implications of these findings are that:

- tumour cell shedding into the blood does not inevitably result in metastasis;
- there is massive redundancy (cell destruction) and selectivity caused by stochastic (chance) factors in the process;
- tumour cell populations which are inconsequential metastatic can still colonise only certain sites, as Paget predicted.

Clinically this information is useful because it provides hope for both physician and patient that the entry of tumour cells into the circulation—for example during surgery, does not inevitably spell doom. It also endorses that knowledge of the patterns of metastatic spread of various tumours is useful for selecting protocols for monitoring recurrence, and it suggests possible new avenues worth exploring for treatment. Subsequent additional work showed that soluble organ specific factors may influence metastatic colonisation and raised the idea that purification and identification of the natural agents which can inhibit formation of metastasis could provide a novel, natural, and possibly non-toxic, form of treatment for established metastatic disease.

The above brief summary indicates that research on tumour invasion and metastasis is becoming very active and is yielding information that is both of potential clinical value and of help for unravelling the mechanisms involved. The pace is accelerating and it would not be unreasonable to expect major advances towards control of this phenomenon in the next few years.

2. Fidler IJ, Kripke ML. Metastasis results from pre-existing variant cells within a malignant tumour. Science 1977;197:893-5.
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