Review

Tumour–stroma interactions
Phenotypic and genetic alterations in mammary stroma: implications for tumour progression
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Abstract

In addition to the well documented role of cytokines in mediating tissue-level interactions, it is now clear that matrix macromolecules fulfil a complementary regulatory function. Data highlighted in the present review extend the repertoire of matrix signalling mechanisms, (1) introducing the concept of ‘matrikines’, these defined as proteinase-generated fragments of matrix macromolecules that display cryptic bioactivities not manifested by the native, full-length form of the molecule, and (2) indicating that a previously identified motogenic factor (migration stimulating factor [MSF]) produced by foetal and cancer patient fibroblasts is a genetically generated truncated isoform of fibronectin, which displays bioactivities cryptic in all previously identified fibronectin isoforms. These observations are discussed in the context of the contribution of a ‘foetal-like’ stroma to the progression of breast cancer.

Keywords: cancer progression, cell motility, epithelial–stromal interactions, fibronectin

Introduction

The pathogenesis of common human cancers is driven by the progressive accumulation of genetic lesions within the target epithelial cell population. Perturbations in the functionality of particular constellations of these oncogenes and tumour suppressor genes gradually confer the emerging population of premalignant and malignant cells with a proliferative advantage compared with their normal counterparts, as well as increasingly aberrant phenotypic characteristics, such as the capacity for local invasion and metastasis [1,2]. During the past few years, emphasis has gradually shifted away from essentially cataloguing these genetic lesions to functional studies that are concerned with their consequences on gene expression and with the intracellular pathways that are responsible for manifesting perturbed epithelial cell behaviour [3].

Although this focus on the target epithelial cell population is central to our understanding of cancer pathogenesis, it has long been our contention that such an exclusively ‘epitheliocentric’ view is too restricted and that regulatory signals that originate in surrounding tissues also make a clinically significant contribution to the kinetics of disease progression [4,5]. This view now sits comfortably within mainstream thought. It is supported by a number of contemporary reviews that concluded that stromal and epithelial cells exert reciprocal effects on the behaviours of each other, and that these essentially epigenetic, tissue-level interactions result in ‘cell activation’ and thereby contribute to tumour progression [6]. No rigorous criteria that define cell activation are currently available. It is generally accepted, however, that these collectively involve the re-expression of certain ‘foetal-like’ phenotypic characteristics, including changes in cell

MSF = migration stimulating factor.
morphology and the secretion of various cytokines, ‘onco-
foetal’ isoforms of matrix macromolecules and matrix-
degrading enzymes. Acting together, these molecules
orchestrate important tissue-level events, including direc-
tional cell migration (chemotaxis), matrix remodelling and the
in-growth of new blood vessels (angiogenesis).

The present short review focuses on the role played by
matrix macromolecules produced by activated cells (both
stromal and epithelial) in the pathogenesis of breast cancer.
Specific attention is given to the following: the manner by
which these molecules co-operate with cytokines in the reg-
ulation of cell behaviour; the expression of relevant cryptic
bioactivities by protease-generated fragments of matrix
macromolecules; recent observations from our laboratory
that indicate that similar cryptic bioactivities are also
expressed by a recently cloned genetically truncated
isoform of fibronectin; and the origins and clinical implica-
tions of stromal cell phenotypic diversity in breast tumours.

Matrix involvement in the mediation of
epithelial–stromal interactions
Interactions between adjacent epithelial and stromal tissues
play a key role in defining the spatial and temporal pattern of
morphogenesis during embryonic/foetal development, and
continue to contribute to the maintenance of normal cell
function throughout adult life [7]. Such tissue-level interac-
tions are mediated by both cytokines and matrix macromole-
cules. A considerable understanding has been gained
regarding the molecular mechanisms by which cytokines
affect cell behaviour. In general terms, these involve the fol-
lowing: cytokine ligation by specific cell surface receptors;
receptor activation and initiation of a chain of intracellular
phosphorylation/dephosphorylation events within diverse
signalling networks; and resultant changes in the pattern of
gene expression and/or functional state of the cytoskeleton.
The inappropriate expression of cytokines and/or their
receptors has long been recognized to contribute to the
pathogenesis of breast cancer [8,9].

In contrast to the well-documented involvement of
cytokines in cancer development, matrix macromolecules
have, until relatively recently, been considered to fulfil an
exclusively structural role that is restricted to providing a
scaffold for cell adhesion and migration. It is now clear,
however, that common matrix constituents elicit signal
transduction cascades as a result of their ligation by inte-
grin receptors, and that such matrix-induced signalling
networks share many common pathways with their
cytokine-induced counterparts [10].

An important biological concept that has emerged during
the past decade is that the bioactivity of cytokines and
matrix macromolecules are mutually interdependent, in the
sense that cytokines and matrix macromolecules exert reci-
procal effects on their respective syntheses (i.e. cytokines
affect the expression of matrix molecules, matrix receptors
and matrix-degrading enzymes, whereas matrix macromol-
ecules modulate the synthesis of cytokines and their
respective cell surface receptors) [11]. In addition, the
effects of cytokines on cell behaviour may be mediated by
the matrix molecules whose synthesis they regulate (i.e.
the observed cellular response to a cytokine may be a
secondary consequence of the bioactivity of the matrix
molecules whose expression is under primary cytokine
control) [11]. Also, matrix molecules bind cytokines and
present them to cell surface receptors in a functionally
optimal state [12]. Finally, matrix macromolecules modu-
late cellular response to cytokines (i.e. the nature of the
macromolecular matrix to which the cells are adherent
determines cellular response to cytokines) [11,13].

With respect to the latter findings, we reported that the
nature of the macromolecular substratum critically deter-
mines whether transforming growth factor-β3 stimulates or
inhibits cell migration and hyaluronan synthesis by dermal
fibroblasts [14]. Taken in conjunction with other published
reports [15], these observations provide the basis of the
‘tissue response unit’ hypothesis [16]. That hypothesis
states that the precise effect of a multifunctional cytokine
on cell behaviour is determined by several tissue-level
parameters, including the presence of other cytokines, the
nature of the macromolecular matrix and the state of target
cell activation. A clear corollary of this model is that the
documented presence of a cytokine during the course of
tumour progression is not sufficient in itself to provide an
insight into its functional significance, in the absence of
complementary data regarding alterations in stroma com-
position and/or physical organization caused by concur-
rent, pathology-associated matrix remodelling.

In addition to the above mechanisms, cell behaviour is also
modulated by the topology of cell–matrix contact. For
example, we have previously reported that vascular
endothelial cells attached to the two-dimensional surface
of a native type I collagen gel continue to express a resting
(‘cobblestone’) phenotype for many weeks when cultured
in the absence of exogenous angiogenic factors [17,18].
We also demonstrated that these cells adopt an angio-
genic (sprouting) phenotype within 24 h of replating within
the three-dimensional gel matrix, again in the complete
absence of exogenous angiogenic factors [17,18]. In this
situation, the only alteration is a shift from conditions in
which the cells contact the collagenous matrix solely along
their basal surface (and can therefore establish an apical-
basal polarity) to culture within an isotropic environment,
in which contact with the matrix is made at sites that are dis-
tributed uniformly along the entire plasma membrane.
Under these conditions, the observed changes in cell phe-
notype, which distinguish resting and angiogenic cells, are
likely to be mediated by topology-dependent alterations in
cytoskeletal organization and functionality [19].
Extrapolating to the in vivo situation, we suggest that the movement of resting endothelial cells from their two-dimensional luminal position into the three-dimensional stromal compartment may be sufficient to induce changes in gene expression that are of relevance to tumour-induced angiogenesis. According to this view, angiogenic factors produced by the tumour may principally function to induce the initial stages of endothelial cell activation and migration into the stroma, and to provide a chemotactic stimulus for the subsequent directed migration of the resultant sprouting cells toward the tumour.

**Matrikines**

An upregulation in the expression of various classes of proteinase has long been recognized to be associated with tumour progression. Certain proteinases (e.g. stromelysin-3) may be described as oncofoetal in that they are constitutively expressed by foetal cells and re-expressed by activated adult cells during the course of tumour progression [20]. A combination of immunolocalization and in situ hybridization studies indicated that these matrix degrading enzymes are commonly coexpressed by both carcinoma and activated stromal cells [21], and available data suggest that the observed anatomical pattern of expression results from reciprocal tumour–stroma interactions [22].

Tumour-associated proteinases have been suggested to contribute to disease progression by a number of means, the most commonly discussed being degradation of basement membrane (thereby facilitating local invasion and metastasis) and release of matrix-bound cytokines [23]. Proteinases may also contribute to cancer progression by degrading matrix molecules into discrete peptide fragments, which display potent bioactivities that are not expressed (i.e. cryptic) by the full-length molecule [23–25]. The ubiquitous matrix macromolecule fibronectin and its proteolytically generated peptide fragments have been particularly well characterized in this regard [26].

**Fibronectin**

Fibronectin is a modular glycoprotein (molecular mass approximately 250 kDa) that consists of a number of protease-resistant functional domains, so named on the basis of their specific binding affinity for other matrix macromolecules and members of the integrin family of matrix receptors (Fig. 1). These domains are as follows: Hep1/Fib1 (low affinity binding to heparin and fibrin), Gel-BD (binding to gelatin/collagen), Cell-BD (binding to integrin receptors on cell surface), Hep2 (high affinity heparin binding) and Fib2 (second fibron binding site). Each functional domain is in turn composed of three possible homology modules (types I, II and III), which contain approximately 45, 60 and 90 amino acids, respectively. Considerably smaller amino acid recognition sequences (such as the RGD tripeptide) function as the minimal peptide motif required for ligation by integrin receptors [27]. All proteolytically generated functional domains of fibronectin, with the notable exception of the 43 kDa gelatin-binding domain (Gel-BD), have been reported to express a diverse range of bioactivities, including effects on cell migration, adhesion, proteinase expression and differentiation [26].

The motogenic activity of full-length fibronectin and its proteolytically generated functional domains have most commonly been studied in variations of the transmembrane (or ‘Boyden chamber’) assay. Previous studies using this assay (for review [28]) have indicated that concentrations of fibronectin and its cell-binding domain in the region of 1–100 µg/ml stimulate the migration of human skin fibroblasts, as well as various other normal and transformed cell types. Ligation of the RGD amino acid motif (which is present in module III-10) by integrin receptors (e.g. \(\alpha_5\beta_1\)) is responsible for mediating motogenic activity.

In view of the important role played by the matrix in modulating cellular response to soluble effector molecules, we developed an alternative migration assay that involves cell attachment and movement within a three-dimensional matrix of native type I collagen fibres. Using this assay, we
reported that purified gelatin-binding domain (Gel-BD) stimulates the migration of human skin fibroblasts, exhibiting a bell-shaped dose–response curve with half-maximal activity manifest at 0.1–1.0 pg/ml (i.e. approximately 2–20 pmol/l) [28]. This unexpected and unprecedented potent motogenic response was further shown to be substratum dependent (i.e. manifested by fibroblasts adherent to a native, but not denatured, type I collagen substratum). The strictly substratum-dependent nature of Gel-BD motogenic activity provides an explanation for the marked discrepancy between results obtained in the collagen gel assay (employing a native collagen substratum) and the majority of published studies using the transmembrane assay (employing denatured collagen-coated membranes).

Subsequent studies indicated that synthetic peptides containing the IGD tripeptide motif (as present with modules I7 and I9 of Gel-BD) essentially mimicked the motogenic activity of Gel-BD, including its substratum dependence [29]. Manifestation of IGD motogenic activity was temporally associated with an increase in the tyrosine phosphorylation of focal adhesion kinase (FAK125) and was inhibited by integrin α5β3 neutralizing antibody [29].

On the basis of these various observations, we suggest the generic term ‘matrikine’ to define proteolytically generated fragments of matrix macromolecules, which display cryptic, cytokine-like bioactivities that are not expressed by their full length precursors. Taken together, these observations reveal a novel functionality of locally produced proteinases and matrix remodelling during tumour progression and other pathologies. They also indicate that, in addition to its other effects on cell behaviour, the extracellular matrix may act as a reservoir of soluble multifunctional bioregulatory molecules.

Migration stimulating factor
We previously reported that foetal skin fibroblasts migrate into gels of native type I collagen to a significantly greater extent than do corresponding adult cells [30]. We also reported that fibroblasts obtained from approximately 50% of sporadic breast cancer patients and more than 90% of patients with familial disease also express a foetal-like migratory phenotype [31–33]. In marked contrast, only 10–15% of fibroblasts obtained from age-matched and sex-matched healthy control individuals in the above studies exhibited a foetal-like pattern of migration. Studies concerned with the biochemical basis of these observations indicated that both foetal fibroblasts and the foetal-like fibroblasts from cancer patients secrete a soluble MSF that is not produced by their normal adult counterparts [34]. Subsequent studies revealed that detectable quantities of MSF were present in the serum of more than 90% of breast cancer patients, compared with only 10–15% age-matched and sex-matched control individuals [35]. Initial biochemical characterization of MSF purified from foetal and cancer patient fibroblast conditioned medium indicated that it had a molecular mass of approximately 70 kDa and contained a peptide sequence identical to a region of the gelatin-binding domain of fibronectin [36,37].

MSF has been cloned from a foetal fibroblast cDNA library (EMBL accession number AJ276395; Schor SL et al., manuscript submitted). Complete sequence data indicate that it is identical to the 5’ end of fibronectin cDNA (including a 56 bp untranslated sequence), with the addition of a novel 175 bp 3’ tail that consists of a 30 bp coding sequence, followed by an untranslated region that contains five in-frame stop codons and a cleavage/polyadenylation signal.

Chromosome mapping and polymerase chain reaction data using template genomic DNA indicate that MSF is a truncated isoform of fibronectin, and that its specific 175 bp 3’ tail is generated by retention of the intron separating fibronectin exons III1a and III1b and its cleavage during subsequent mRNA maturation. The deduced MSF protein consists of the intact 70 kDa amino-terminus of fibronectin (containing the entire Fib1/Hep1 and Gel-BD regions, and the first portion of module III1); and a unique 10 amino acid carboxyl-terminus (VSIPPRNLGY), as coded by the first 30 bp of the retained intron (Fig. 1). All previously identified fibronectin isoforms (of which there are approximately 20) are ‘full-length’ compared with MSF and are produced by well characterized alternative splicing events at three sites downstream from the newly identified MSF splice site (i.e. the exons coding for the ED-A and ED-B type III homology modules, as well as the IIICS region; Fig. 1).

Recombinantly expressed human MSF stimulates the migration of adult skin fibroblasts with a dose-response relationship that is identical to that of Gel-BD [25] (Schor SL et al., unpublished data). Significantly, MSF motogenic activity is completely abolished by an anti-Gel-BD monoclonal antibody and is unaffected by antibodies raised against all other fibronectin functional domains, including the amino-terminal Fib1/Hep1 domain. Taken together, these (and related observations) suggest that MSF motogenic activity is mediated by a peptide region that is located within its gelatin-binding domain; and that the IGD motif is a prime candidate bioactive sequence. As is the case with Gel-BD, fibroblast motogenic response to recombinant human MSF is strictly matrix dependent, being manifest by cells that are adherent to native, but not denatured, type I collagen. Recombinant human MSF and Gel-BD both express a number of other relevant bioactivities, including the stimulation of hyaluronic acid synthesis [38] and the induction of a potent angiogenic response in the chick yolk sac membrane (Schor AM, unpublished data).

On the basis of these various observations, we conclude that MSF and Gel-BD are functionally equivalent, but pro-
duced by distinct mechanisms, presumably under independent control (i.e. Gel-BD is a matrikine that is generated by the proteolytic degradation of matrix fibronectin, whereas MSF is the first genetically truncated isoform of fibronectin identified). It should also be noted that MSF differs from all fragments of full-length fibronectin isoforms (including Gel-BD) in terms of its unique 10 amino acid carboxyl-terminus. The generation of MSF by a genetic mechanism therefore affords the cell with a means to selectively produce a functional equivalent to Gel-BD without the necessity of producing a mixture of proteinase generated matrix fragments that express a complex array of cryptic bioactivities.

Using an MSF-specific antisense riboprobe (to a sequence contained within its unique 175 bp 3′ tail), in situ hybridization revealed that MSF is weakly expressed by a small number of ductal epithelial cells and interlobular fibroblasts in normal breast, and is more strongly expressed by carcinoma cells, stromal fibroblasts and some blood vessels in breast tumours (Schor SL, unpublished data). These observations are consistent with data from our laboratory that indicate that MSF is constitutively expressed by interlobular, but not intralobular fibroblasts isolated from normal breast tissue [39]. Thus, they indicate for the first time that MSF is abundantly and concurrently expressed by breast carcinoma cells, as well as tumour-associated fibroblasts and vascular endothelial cells. Complementary immunolocalization using an MSF-specific monoclonal antibody confirmed the association of MSF protein with both mammary carcinoma and stromal cells. The expression of MSF by both carcinoma and tumour-associated stromal cells is consistent with published data that indicate a similar concurrent expression of full-length oncofoetal fibronectin isoforms by these cells [40]. A functional co-operation between MSF and previously described oncofoetal fibronectins in the induction of new blood vessels is suggested by the angiogenic activity of MSF and the reported expression of oncofoetal fibronectins that contain the ED-A and ED-B domains in vascular-rich stroma [41,42].

It should finally be noted that the presence of MSF in tumour tissues (as indicated by both immunolocalization and in situ hybridization) is not in itself sufficient to conclude that MSF bioactivity is expressed. In accord with the tissue response unit hypothesis, the effect of modulating parameters, such as the nature of the extracellular matrix and cytokine profile, must also be taken into account. With respect to the latter, we previously reported that transforming growth factor-β1 is a potent inhibitor of MSF and Gel-BD functionality [28,43]. Our current understanding of MSF expression and functionality during breast cancer progression is summarized in Fig. 2.

Figure 2

The expression and functionality of migration stimulating factor (MSF) during the progression of breast cancer. MSF is concurrently expressed by a proportion of carcinoma cells, stromal fibroblasts and vascular endothelial cells (producer cells). In vitro and in vivo observations indicate that it affects a diverse range of phenotypic features of these same cells (now functioning as target cells). MSF activity may be mediated by autocrine mechanisms (i.e. same producer and target cell population) and paracrine mechanisms (i.e. different producer and target cell populations). Both MSF expression and cellular response to it are matrix dependent: HA, hyaluronic acid.

Origins of stromal cell phenotypic diversity

We originally speculated that ‘foetal-like’ fibroblasts were associated with breast and other types of cancer, and that the resultant perturbations in signalling between stroma and epithelium played an important role in driving the process of tumour progression [4,5]. In these models, fibroblasts were defined as ‘foetal-like’ on the basis of their re-expression of molecules (such as MSF) that are constitutively expressed during development. Subsequent studies considerably extended these analyses to include many other gene products [44,45] and explicitly went on to support the view that stromal ‘foetalization’ is an important feature of carcinogenesis [46].

These variations in stromal cell phenotype may arise through both genetic and epigenetic mechanisms. With
respect to the former, Moinfar et al. [47] recently presented evidence documenting loss of heterozygosity in DNA isolated from microdissected regions of mammary tumour stroma that were distinct from those associated with the carcinoma cells. These observations support the view that environmental carcinogens that were previously implicated in the development of breast and other cancers also produce genetic lesions in stromal cells, and that this genetic damage may also contribute to the course of disease progression. Alterations in stromal cell phenotype may additionally arise from epigenetic mechanisms that involve responses to altered signalling from a progressively aberrant epithelium and to stress-inducing micro-environmental agents (as contained in cigarette smoke). In this context, we recently observed that a number of cytokines and genotoxic agents ‘switch on’ MSF expression by adult skin fibroblasts and that, under certain circumstances, this alteration in gene expression is persistent for many months in vitro (Schor SL et al., unpublished data).

Conclusion

An important function of stromal cells is the deposition and remodelling of the extracellular matrix. In addition to providing the physical basis for tissue integrity, it is now clear that matrix macromolecules exert profound effects on cell behaviour by a variety of mechanisms. These include the following: receptor-mediated signal transduction; modulation of cellular response to cytokines; binding and presenting cytokines in an optimal conformation; and acting as a reservoir for proteinase-generated matrixines. Truncated isoforms of matrix molecules produced by alternative splicing (such as MSF) may also express a range of bioactivities that are cryptic within the full-length molecule. The substratum dependence of Gel-BD and MSF serve to emphasize the critical importance of the physical organization of the matrix in terms of its effect on the behaviour of adherent cells.

We previously speculated that expression of MSF and other oncofoetal effector molecules by the foetal-like (activated) stromal cells in cancer patients may contribute to disease progression by creating a milieu that promotes the clonal expansion and manifestation of invasive behaviour of the emerging (pre)malignant cell population [5]. In this regard, it is important to bear in mind that tumour progression is a relatively indolent process, in which many decades may elapse between inception of the initiating genetic lesion and the emergence of a clinically recognizable malignancy. Available data suggest that the proportion of ‘initiated’ cells that eventually develop into a tumour is quite low. We accordingly suggest that factors that may alter the kinetics of progression may play an important, and perhaps decisive, role in determining the probability of disease inception. It is in this postulated role of an ‘accelerator’ of cancer progression that we envision the contribution of activated ‘foetal-like’ cells (and the MSF they produce) to cancer pathogenesis. Seen in this light, MSF functions as a severity (rather than that causative) agent. The documented bioactivities of MSF (motogenic, angiogenic and stimulation of hyaluronan synthesis) are all potentially relevant features of its functionality. In the light of these findings, we suggest that the development of adjunct therapies that are specifically designed to normalize activated cell function may prove clinically efficacious in the treatment of patients with cancer and/or the retardation of cancer development in healthy individuals who are deemed to be at elevated risk.

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