‘Malignant melanoma microecosystem’: Immunohistopathological insights into the stromal cell phenotype (Review)

PASCALE QUATRESOOG, MARIE-ANNICK REGINSTER and GÉRALD E. PIÉRARD

Department of Dermatopathology, University Hospital of Liège, BE-4000 Liège, Belgium

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Abstract. Cutaneous malignant melanoma (MM) is rooted in the dermal connective tissue, which consists of apparently unremarkable stromal cells as they appear upon regular histopathological examination. However, a number of in vitro studies have shown that these cells produce diverse types of cytokines, growth factors and enzymes in excess. In addition, they store and probably release various structural components of the extracellular matrix (ECM). Most of the current information comes from in vitro experiments, and these findings do not always correlate with investigations carried out using excised human MM tissue. The MM-stroma connection appears crucial to the regulation of neoplastic growth, invasiveness and initial metastatic spread. However, little is known about the in vivo intracellular storage and extracellular deposits of specific ECM macromolecules located inside and around MM lesions. This review summarizes various distinct features of the peri-MM stroma, which shows an intracytoplasmic abundance of Factor XIIIa, versican and various α (IV) collagen chains. The area exhibiting such changes corresponds to the location where neoangiogenesis commonly develops and where extravascular unicellular metastatic MM lesions are possibly found. Some of these inconspicuous migratory malignant melanocytes may actually correspond to MM stem cells. Their presence was found to be significantly associated with an increased risk for distant metastases, particularly in the sentinel lymph nodes. Although much remains to be learned, active intervention of the ECM appears likely in the inconspicuous early dermal metastatic migration of MM cells.

Contents

1. Introduction
2. Materials and methods
3. Malignant melanoma micrometastases
4. Stroma immunohistochemistry beneath malignant melanoma
5. Discussion
6. Conclusion

1. Introduction

Cutaneous malignant melanoma (MM) is basically an uncontrolled overgrowth of neoplastic melanocytes. At some stage of its progression, the neoplasm exhibits a high metastatic potential. It proves to be resistant to drug-induced apoptosis, which is believed to underlie the resistance of MM to conventional chemotherapy and radiotherapy (1,2). Various interactions exist between MM cells and other biological systems, including immune cells, vascularity, contiguous stromal cells and the dermal extracellular matrix (ECM). Certain aspects of MM-stroma interactions are thought to be associated with disease prognosis (3). In addition, environmental influences, including ultraviolet (UV) light, are probably responsible for MM initiation and may support its progression along with the intervention of diverse autocrine and paracrine factors (4). In particular, a number of growth factors and specific enzymes are released in the MM microenvironment (5-7).

The participation of the host in the ‘cancer microecosystem’ basically involves the microvasculature, stromal cells and specific immune reactions (8-10). Angiogenesis is a typical host-mediated response to many cancers. It appears crucial for cancer progression, as blood vessels deliver nutrients and oxygen to neoplastic cells (11). Furthermore, the microvasculature likely allows communication between the primary MM and its metastases. Pro-angiogenic molecules originate from cancer cells as well as from the stroma. The relative contribution of both compartments is likely to change with MM type and site, and is balanced by other factors as well (11,12).

Cross-talk between MM and stromal cells may be mediated through direct heterotypic cell-cell contacts, adhesion molecules, signaling factors, and other secreted molecules consisting of growth factors, cytokines, chemokines, ECM proteins, proteinases, proteinase inhibitors and lipid products (13). Conceptually, the MM microenvironment is crucial for the maintenance of cellular functions and tissue integrity, suggesting that a cancer-induced change in the ECM may contribute to cancer invasion (14). Any alteration in the MM stroma may be due to an imbalance in the cytokine profile, resulting from oncogenic changes in the cancer cells. In
particular, experimental animal models have demonstrated that cancer invasion is stimulated by the wound-healing stroma (15).

Both stromal cells and the ECM located beneath primary MM lesions are therefore likely involved in the process of invasion of the neoplasm and in the early dissemination of micrometastases associated or not with neoangiogenesis (2,10,16-19). These characteristics are possibly associated with phenotypic changes in the stromal cells in the MM vicinity. In recent years, tumor growth regulation by ECM components has been one of the main topics of neoplastic biology research.

2. Materials and methods

This study is a review of current peer-reviewed publications admixed with personal original findings from a series of 400 MM cases with a thickness ranging between 0.4 and 1.0 mm (median 0.83) that were retrieved from our files. The microscopic diagnosis was previously established by a group of three dermatopathologists. Immunohistochemistry was performed as previously described (20-23). In short, samples were fixed in buffered formalin and embedded in paraffin. A series of 6-µm sections were prepared for immunohistochemistry. The avidin-biotin peroxidase method was used with the antibodies listed in Table I. After 1 h of incubation with each primary antibody, the slides were washed in Tris-buffered saline (TBS) and incubated for 30 min with the secondary antibody (biotinylated swine anti-rabbit, 1:300; Dakopatts Glostrup, Denmark). Slides were rinsed in TBS and covered by the EnVision (Dakopatts) polymer-based revelation system. After TBS washings, Fast Red (Dakopatts) was used as the chromogen substrate. The final steps consisted of counterstaining with Mayer’s hemalum and mounting in glycerin mounting medium (Dakopatts). Negative immunohistochemical controls were performed by omitting or substituting the primary and the secondary antibodies from the laboratory procedure.

3. Malignant melanoma micrometastases

A typical biological feature of human MM is the tremendous impact of the primary lesion thickness on prognosis. Primary MM <1 mm in thickness is associated with a high cure rate, sharply contrasting with thicker lesions associated with poorer prognosis. The apparent breakpoint beyond an ~1-mm thickness is a discouraging factor in disease outcome. One possible reason appears to be linked to vascularization patterns of MM (11,12). Such an anatomic argument is persuasive, but it is by no means the only one.

MM cells apparently fail to form metastases unless they present the genotypic and phenotypic information allowing them to effectively migrate in the ECM, intravasate, extravasate, cross interstitial basement membranes and proliferate in distant tissue sites. These characteristics are expressed by variant subpopulations of metastatically competent MM cells present in primary neoplasms (2). These subpopulations probably acquire a growth advantage at the primary site over time, so that they become a dominant proliferating population. At this stage, the MM truly expresses overt malignancy. As a result of this process of clonal dominance of metastatically competent cells, it is possible that most thin primary MM lesions contain very few, if any, metastatically competent cells, whereas thicker MM lesions may contain significant proportions of such cells. It is possible that the stromal microenvironment plays a role in such a shift in the biological profile of MM (24).

<table>
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<th>Table I. Panel of antibodies.</th>
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<tr>
<td><strong>Antigen</strong></td>
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<tr>
<td>α1 (IV) collagen</td>
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<td>α5 (IV) collagen</td>
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<tr>
<td>Elafin</td>
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<td>Factor XIIIa</td>
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<td>Lysozyme</td>
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The four metastatic routes. In order to form metastases at distant sites, MM cells must acquire certain functions and properties in an ordered sequence referred to as the MM metastatic cascade. In order to form overt metastasis, this process must encompass hyperproliferation, detachment from the primary neoplasm, invasion into the peritumoral stroma and possibly penetration into blood and lymphatic vessels, survival in the circulation, adhesion to a vessel wall at the site of the final metastatic deposition, extravasation and proliferation (3).

The specific function involved in the metastatic cascade combines intrinsic characteristics of various MM cells and regulatory influences from the microenvironment. Indeed, MM cells and their surrounding stroma jointly form a microecosystem receptive or not to early conspicuous metastatic spread.

Early MM micrometastases are not discernable upon regular clinical or dermoscopic examination. They are disclosed under the microscope, particularly after highlighting their presence using immunohistochemistry (2,17). They are found in four distinct locations, namely i) inside lymph vessels, ii) inside blood vessels, iii) in a perivascular location just adjacent to the outer area of the endothelial lining; and iv) dispersed inside the stroma (10,16,18). The latter eventuality is not infrequently associated with neoangiogenesis, and an enhanced neoplastic germinative pool is commonly found (10,19).

The active migration of metastatic MM cells in the peritumoral stroma is probably a complex process. It involves the active mobility of MM cells and changes in the neoplastic cell adherence systems with ECM components.

Melanoma stem cells. The presence of MM stem cells is an important consideration when investigating the characteristics of MM micrometastases and their relationship with the peritumoral stroma (10,22,25-27). Similar to physiological stem cells, cancer stem cells are capable of self-renewal and differentiation, and have the potential for indefinite proliferation, a function linked to MM growth (2,22,28). Although conventional anti-cancer treatments may eradicate most malignant cells, they are potentially ineffective against chemoresistant cancer stem cells, which are ultimately responsible for tumor recurrence and
progression (2,10,25). MM shows tumor heterogeneity, undifferentiated molecular signatures and increased tumorigenicity of MM subsets with embryonic-like differentiation plasticity. This strongly suggests the presence and involvement of MM stem cells in the initiation and propagation of this malignancy (25-27,29-32). The ECM structure and biologic activity may influence the invasiveness and propagation of MM stem cells.

**Micrometastases and the peri-melanoma stroma.** When present, interstitial unicellular MM micrometastases are frequently found and confined to the perineoplastic stroma. Their presence is significantly correlated with the risk of involvement of the sentinel lymph node (17).

**4. Stroma immunohistochemistry beneath malignant melanoma**

Upon standard histopathological examination, stromal cells appear normal underneath primary MM lesions when partial regression is not operative. However, their differentiation as revealed by immunohistochemistry appears altered when compared to the surrounding skin. In particular, phenotypic changes are noted when identifying the transglutaminase Factor XIIIa, α (IV) collagen chains, as well as elafin and versican. It is possible, although not yet proven, that transforming growth factor (TGF)-β1 and platelet-derived growth factor (PDGF) may play a role in the alteration of the stromal host compartment in MM.

**Factor XIIIa-enriched stromal cells.** Factor XIIIa-enriched stromal cells are commonly identified as dermal dendrocytes (DDs). They are preferentially found adjacent to superficial microvasculature (24,33-35). Increased numbers of Factor XIIIa-positive DDs are often found in the vicinity of most invasive cutaneous neoplasms. In our experience, Factor XIIIa-positive DDs are numerous; they neighbor and infiltrate most thin MM lesions (24). By contrast, they are present in few numbers or even absent in thick primary MM lesions and their metastases (36-38). Circumstantial evidence indicates that the density of Factor XIIIa-positive DDs is correlated with a low proliferative rate of MM cells. Thus, Factor XIIIa-positive DDs may not be passive bystanders in MM (24,34,36). Their function may differ based on whether they are located in the stroma or inside the neoplasm (24). Intratumoral DDs may be associated with a growth-restricting role. By contrast, stromal DDs may help in the invasiveness and metastatic spread of MM cells.

**Collagen IV-enriched stromal cells.** In malignant neoplasms, basement membranes (BMs) are composite structures synthesized by tumor cells or stromal cells; either by one of these two cell types yet dependent on the interactions between them, or a mixture from both origins. These tumoral BMs are often abnormal in their composition and ultrastructural features (39,40). BM material appears to accompany malignant cells rather than to prevent invasion as a physical barrier. Nevertheless, active interactions between neoplastic cells and stroma, in particular the ECM, play a key role in neoplastic progression leading to invasion and metastasis (41). Several BM components have been identified surrounding MM cells, including collagen IV (39,42-44). In the skin, collagen IV represents an assembly of α1 (IV) and α5 (IV) collagen chains. In MM, some neoplastic and stromal cells exhibit intracytoplasmic immunolabeling for α1 (IV) chains (44). The pattern is heterogeneous. BM components, including collagen IV, gradually disappear during the dermal ingrowth of MM cells. Notably, a minority of MM cases without any identifiable micrometastasis and a majority of MM with cutaneous micrometastasis show discrete cytoplasmic positivity for the α5 (IV) collagen chain (44).

Distribution of the α1 (IV) collagen chain in MM highlights the heterogeneity in both cell differentiation and stroma-MM interactions. Thus, MM cells appear to have their own individual potential to be enclosed by a BM and to interact with the stroma. This biological aspect may be related to neoplastic progression and may influence inconspicuous metastatic potential.

**Versican-enriched stromal cells.** Versican is a large proteoglycan normally present inside the stromal cells of the skin. The molecule belongs to the chondroitin sulfate family of the hyalectan group, named for its ability to bind hyaluronan (45). In mammals, versican appears as four possible spliced isoforms, V0 to V3. Little is known concerning the differential regulation of the isoforms or about their respective roles in
the ECM either normal or peritumoral. Versican production is deregulated in several types of human cancer (46). As it is largely expressed in rapidly growing neoplastic cells, it has been suggested that versican plays a direct role in cell proliferation and other cell functions (45). It appears to be particularly abundant in the stromal cell population underlying MM (Fig. 1) (46-49). Versican overexpression was found to sharply circumscibe to a cup-shaped structure cuffing the bottom of MM lesions. In addition, some nests of MM cells were found to be strongly labeled with the anti- versican antibody (Fig. 2). This finding contrasts with another study reporting the absence of versican immunoreactivity in neoplastic melanocytes (49). In addition, versican expression is not correlated with Breslow tumor thickness and Clark’s level (49).

Elafin-, versican- and lysozyme-loaded ECM. Elastic fibres are coated with distinct molecules following chronic UV exposure (50-53). The serine anti-leukoprotease elafin, as well as versican and lysozyme, bind to elastin preventing elastolytic degradation by elastases on sun-exposed areas exhibiting solar elastosis (51,53,54). Under these conditions, the labeling was found to range from partial, moderate to strong. In addition, inhibition of elastase may decrease the adhesion of cancer cells to endothelial cells (55). In addition, elafin was reported to elicit p53-dependent apoptosis in cultured MM cells transfected by a plasmid-producing elafin under doxycycline boosting (56). In contrast to these in vitro experiments, immunohistochemistry did not disclose an intratumoral cell presence of elafin in human MM. Rather, keratinocytes covering MM overexpressed elafin in their cytoplasm. Of note, Western blotting and reverse transcription analyses indicated transcriptional elafin repression in MM cells (56).

The implication of elafin in other diseases, such as psoriasis and graft-versus-host reaction, indicates its distinct importance in skin biology (57,58).

5. Discussion

This review highlights the existence of a distinct region of the dermis adjacent to the base of a primary MM lesion. Stromal cells exhibit particular phenotypic features suggesting altered functionality. The involved territory appears to be conducive to micrometastatic spread. Some of these cells survive singly, and due to their manner of migration to other organs may represent MM stem cells.

In addition to the importance of MM vascularization for tumor growth, invasiveness and metastatic spread (10-12,19,59), numerous other roles are ascribed to the tumoral stroma. This structure is involved in a constant remodeling following degradation and repair of the ECM. Notably, immunohistochemistry highlights the direct implication of MM cells in the synthesis and/or storage of certain ECM molecular components.

The immunohistochemical characterization of MM cells is important (20,22,60-64), yet should be extended to the peritumoral stroma, including the microvasculature (10-12,59) and other ECM components. A comprehensive mapping of MM immunohistochemical characteristics should aid in identifying relevant targeted therapies (63-66). Inflammatory cells and immunocytes represent another class of host cells that are regulated by the balance of cytokines. They perform counter-current invasion, from the circulation into the tumor, and provide routes for MM cell invasion. It is important for our understanding of MM stroma turnover that tumor-infiltrating leukocytes produce proteases.

6. Conclusion

Interaction between MM and its stroma is evident during the invasive and metastatic stages of disease progression. Stromal cells are known to secrete metalloproteinases and their inhibitors, growth factors, the scatter factor/hepatocyte growth factor and other factors, as well as participate in the growth and mobility of MM cells. In addition, other molecules are synthesized and overexpressed by stromal cells and/or MM cells. Immunohistochemistry has identified Factor XIII-a, α1 and α5 (IV) collagen chains, versican, elafin and lysozyme. These possibly influence the migration of MM cells, including their stem cells.

While MM cell motility cannot be directly assessed, there is circumstantial evidence indicating that motility is essential to MM progression and possibly of prognostic significance. Apart from the secretion and activation of enzymes altering the ECM, a variety of stromal alterations occur following overexpression of diverse ECM components. Molecular morphology yields evidence suggesting that the MM stroma plays an integral role in MM. Although much remains to be determined, the findings as described in the present review may have diagnostic and prognostic significance, which warrant further investigation.

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