Extracellular Matrix Macromolecules in Tumour Microenvironment with Special Reference to Desmoplastic Reaction and the Role of Matrix Proteoglycans and Hyaluronan

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Abstract

Cancer cells create their own microenvironment via dynamically interacting with the surrounding non-malignant stromal cells and various Extracellular Matrix (ECM) macromolecules as well as with a large number of other proteins within the ECM.

The term desmoplastic reaction is used to describe a phenomenon in which ECM macromolecules including specific species of proteoglycans and hyaluronan variously accumulate around the tumour mass. The precise role of the desmoplastic reaction is not known. However, it has been proposed to represent either a defense mechanism by non-malignant host cells against invading cancer cells or to be a tumourigenesis promoting phenomenon induced by cancer cells to assist their propagation. Even an idea that desmoplastic reaction precedes the development of cancer has been introduced, further emphasizing the importance of ECM macromolecules in tumourigenesis. Here, we will discuss ECM macromolecules in tumour microenvironment focusing particularly on desmoplastic reaction and ECM proteoglycans and hyaluronan. We will also present examples of strategies how ECM macromolecules involved in desmoplastic reaction could be considered in the development of innovative oncolgical pharmacotherapies in the future.

Keywords: Extracellular matrix; Tumour microenvironment; Desmoplastic reaction; Cancer; Proteoglycan; Decorin; Versican; Hyaluronan


The Extracellular Matrix of Tumour Microenvironment

In general, there is probably no disease without changes in the composition of the ECM. The macromolecules of the ECM form a complex 3D network composed of collagens, elastin, glycoproteins, Proteoglycans (PGs) and Hyaluronan (HA) [1]. Furthermore, in the ECM there are versatile groups of additional molecules such as ECM -modifying enzymes, ECM-binding growth factors and other ECM-associated proteins [2]. The overall composition and structure of the ECM are unique and highly dynamic in each organ and tissue. Molecules of the ECM provide not only structural support but they also crucially regulate cell adhesion, migration, proliferation, differentiation, and survival [1-3]. In normal situation, molecules of the ECM control the behaviour of cells through feedback loops maintaining tissue homeostasis [1,2]. In tumours, on the other hand, there is their own microenvironment where ECM molecules variously promote tumour cell survival, growth, migration and metastasis [4]. Within the ECM there is also a rich cellular component consisting of normal fibroblasts, activated fibroblasts called myofibroblasts or cancer-associated fibroblasts, inflammatory cells and tumourigenesis promoting stem cells or their derivatives [5,6]. The complex interplay between the cancer cells, non-malignant host cells and various ECM molecules leads to vast activation process resulting in desmoplastic reaction which means the accumulation of ECM macromolecules such as collagens, matricellular proteins, proteinases and their inhibitors and also specific species of PGs and HA.

Desmoplastic Reaction in Cancer

The composition of tumour associated desmoplastic reaction is very similar to that of the granulation tissue [7,8]. The term desmoplastic reaction describes a phenomenon in which a dense fibrotic tissue rich in collagen and other ECM macromolecules including specific species of PGs and HA surrounding the tumour mass is observed [9,10]. This reactive stromal response is thought to

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Received April 02, 2013; Accepted April 26, 2013; Published May 13, 2013


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be produced primarily by specified myofibroblasts which form a highly heterogeneous cell population with different phenotypes and modulate the behaviour of both tumour cells and host cells [6,8]. The exact role of the desmoplastic reaction associated with cancer development is not known. However, it has been proposed to represent either a defense mechanism by normal host cells against invading cancer cells [11,12] or to be a tumourigenesis promoting phenomenon induced by cancer cells to assist their propagation [7,13]. Furthermore, an idea that desmoplastic reaction precedes the development of cancer has been introduced [8,14]. Nevertheless, the desmoplastic reaction is clearly detectable in many primary and metastatic tumours including colorectal [12], liver [15], pancreatic [10,16] and prostate cancers [12] and even in some rare tumour types such as syringomatous carcinoma [17]. However, further studies are needed to better clarify the exact role of desmoplastic reaction in tumourigenesis before it is possible to fully assess whether desmoplastic reaction associated molecules might provide a target in the treatment of cancers.

**Extracellular Matrix Macromolecules Associated with Desmoplastic Reaction**

In the ECM, there are several individual macromolecules that have been shown to be associated with desmoplastic reaction. These molecules have various effects on cancer cells depending on both the molecule and the cancer type in which they are expressed. Overexpression of collagen types I, III, IV, V and XII create a sclerotic stroma around the cancer cell population comprising a dense fibrillar deposition typical of desmoplastic reaction [12,18]. In addition to this restrictive fibrillar formation, different collagen types can also have distinct functional roles. For example, the accumulation of collagen type I in pancreatic cancer has been shown to increase the malignancy of the cancer cells via increasing their proliferation, migration and survival [19,20]. More precisely, in pancreatic cancer it has been shown that cancer cells can increase the expression of collagen synthesis by normal pancreatic stellate cells thus providing a functional interaction between the malignant and normal cells [19]. The collagen-rich tumour environment has also been demonstrated to have a biochemical role affecting the cancer progression through the interplay between type I collagen, membrane-type I matrix metalloproteinase (MT1-MMP) and Transforming Growth Factor Beta (TGF-β) [20]. In the case of type IV collagen, its upregulation has been demonstrated to be associated with enhanced metastasis power of colorectal cancer cells to the liver [21].

Besides collagens, matricellular proteins such as thrombospondin-1 (TSP-1), -2 (TSP-2) and tenasin-C (TNC) can have altered expression patterns in different cancers [22,23]. Contrary to type I collagen, overexpression of TSP-1 has been suggested to have a protective effect on tumour progression, particularly in papillary thyroid carcinoma where the interaction of TSP-1 with one of its receptor, alpha root of beta 3 is of central importance [24]. On the other hand, tumour-derived pancreatic stellate cells can promote the release of growth factors such as TSP-2 which has been shown to stimulate pancreatic cancer cell invasion [25]. Regarding TNC, its increased expression has been shown to be associated with tumour aggressiveness in medullary thyroid carcinoma [26]. In addition to collagen and matricellular proteins, also proteinases as well as their inhibitors can contribute to desmoplastic reaction.

Especially, the presence of Matrix Metalloproteinases (MMPs) and their inhibitors, namely Tissue Inhibitors Of Metalloproteinases (TIMPs) are of central importance when predicting the invasion and metastasis of cancer cells [20,27]. In breast cancer, it has been shown that the expression of both MMPs and TIMPs is significantly associated with the capability of tumour cells to metastasize [28,29].

Although TIMPs are usually considered to act as inhibitors and regulators of MMPs, they have also been demonstrated to possess distinct tumourigenesis promoting functions, thus further emphasizing their multifunctionality [29,30]. All the aforementioned molecules associated with desmoplastic reaction emphasize the vital interaction between the stroma and the cancer cells. This is also true for ECM PGs and HA introduced and discussed next.

**Extracellular Matrix Proteoglycans and their Classification**

Proteoglycans are essential macromolecules composed of a specific core protein substituted with covalently linked Glycosaminoglycans (GAGs) side chains, the hallmark of the PGs. HA is an exception; it is the only GAG synthesized in a free form lacking a protein core. GAGs are linear, negatively charged polysaccharides which exist as sulfated (Chondroitin Sulfate (CS), Dermatan Sulfate (DS), Keratan Sulfate (KS), heparin, Heparan Sulfate (HS)) and non-sulfated forms and together with the core protein create the physiological properties of each individual PG [31].

PGs such as decorin and biglycan are found in almost all extracellular matrices of tissues but some PGs seem to be highly tissue and cell specific [32]. Together with various combinations of core proteins and different GAGs, PGs have a vast diversity of biological functions.

Extracellular PGs can be classified in several ways. On the basis of their localization, three main groups of ECM PGs exist, namely matrix-accumulated PGs, Small Leucine-Rich PGs (SLRPs) and basement membrane PGs [33]. When taking into account also the size and modular composition of ECM PGs, in addition to the localization and the type of GAG carried by the PG’s core protein, division to the following families can be made: SLRPs, modular PGs, and cell-surface PGs [31,32]. As the knowledge of the physiological functions and the pathological roles of PGs expand in time, PG classification will modify itself further.

The expression of PGs and HA is known to be significantly different in tumour tissues compared to normal ones [12,34]. In tumour tissues, the unique expression of these ECM molecules can variously modulate tumourigenesis, for example by regulating growth factor activity and thereby tumour cell proliferation and invasion [35]. Next, we will discuss the ECM PGs and HA in cancer associated desmoplastic reaction (Table 1) and review their potential roles in tumourigenesis.

**SLRPs in tumour microenvironment**

Small Leucine-Rich Proteoglycans (SLRPs) such as Decorin (DCN) and biglycan (BGN) are secreted PGs composed of a protein core with Leucine-Rich Repeats (LRRs) and at least one GAG side chain [32,36]. Both DCN and BGN contain 10 LRRs and comprise the first sub-family of SLRPs. Altogether there are five sub-families (Class I-V) with 18 known members [31,36]. The structure of SLRPs enables them to modulate cellular functions through interactions with other ECM proteins, various cytokines, cell surface receptors and different growth factors [37]. In addition to general effects such as modulation of adhesion, proliferation and migration, SLRPs possess also molecule-specific functions which are achieved in a cell-specific manner [31].

Although the role of SLRPs in desmoplastic reaction is not exactly
known, several of them are associated with cancer development. Regarding decorin, its expression has been shown to vary in cancers. For example, in Kaposi’s sarcoma DCN expression is markedly upregulated around the tumour mass (Figure 1) [38]. In contrast, in colon carcinoma the expression of DCN is reduced [39]. However, the current view is that DCN possesses antitumourigenic and antimitastatic functions via various mechanisms of action [40-43]. DCN has a recognized role in the reduction of cancer cell invasion and migration even as an aberrantly expressed nuclear localized form [44]. Specifically, DCN inhibits tumour growth by antagonizing various receptor tyrosine kinases such as Epidermal Growth Factor Receptor (EGFR) and other ErbB family members resulting in cell cycle arrest through increase in endogenous p21 [45-48]. Furthermore, analogously to EGFR signaling, DCN’s interaction with mesenchymal-epithelial transition (Met) receptor, i.e., receptor for hepatocyte growth factor has been shown to block signaling pathways leading to e.g. the inhibition of angiogenesis [48-51]. Decorin also sequesters multiple other growth factors 51 , particularly Transforming Growth Factor 1 (TGF–1), and thereby promotes tumour progression [51-53]. The role of DCN in tumourigenesis can also be mediated via the Insulin-Like Growth Factor-1 (IGF–1) system [54]. In addition, although not well understood, DCN’s interaction with Toll-Like Receptors (TLRs) has been reported to cause reduced tumour growth via immune response through Programmed Cell Death 4 (PDCD4) and microRNA-21 [55].

Antitumouric action of DCN has been observed with systemic delivery of decorin core protein [56] and with DCN transfection on cancer cells [57,58].

The association of decorin’s class mate, BGN, with cancer is more complicated. In several cancer types such as gastric [39] and colorectal cancers [60] decreased expression of BGN seems to be a biomarker for poorer prognosis. Furthermore, in pancreatic adenocarcinoma, increased expression of BGN is considered to prevent tumour growth and invasivity of cancer cells [61]. BGN has also been discovered to act as a signaling molecule [62,63] and being capable of regulating downstream signaling events by clustering different types of cell surface receptors [64]. While BGN mediated signaling influences tumour microenvironment, the precise role of BGN in cancer is still not known. In a recent paper by Yamamoto and colleagues [65], BGN was found to act as an autocrine angiogenic factor of tumour endothelial cells, thus widening the role of BGN in tumourigenesis even further.

Also other SLRPs including asporin [66,67], fibromodulin [12,17,68], and lumican [69-71] have been shown to be associated with tumourigenesis - asporin and fibromodulin via their potential clinical value for applications in diagnostics and therapeutics and lumican via its antiangiogenic activity. In more detail, asporin is found to be expressed in significant amounts in pancreatic ductal adenocarcinoma tissue [66] and its concentration has been shown to be increased in the blood of men with prostate adenocarcinoma [67]. Regarding fibromodulin, it exhibits exclusive ectopic expression in B-cell chronic lymphocytic leukemia [72]. Lumican, in turn, has been shown to enhance Fas mediated endothelial cell apoptosis in vivo [71] as well as cause interference with 1 receptor activity and downregulation of tumour microenvironment.

Expression of modular PGs and HA in tumour microenvironment

Modular ECM PGs are multidomain PGs with various elongated protein modules and highly glycosylated structure [32]. They can be divided into two groups; first to hyalectans such as versican, aggrecan and brevican, and secondly to non-hyalectans binding PGs, e.g. perlecan [32,73]. These ECM PGs contain a central domain that carries most of the GAGs, from three in brevican up to 100 in aggrecan, and this domain is flanked by regions that bind HA and contain C-type lectin-like domains, respectively [32,74]. From the hyalectans, versican is the most cancer associated ECM PG, as its expression is altered in most cancer types including bladder [75], colon [76], breast [77] and ovarian cancers [78]. In adenocarcinomas, versican seems to be the primary ECM PG secreted by the activated peritumoural stromal cells, e.g. myofibroblasts [79,80]. The role of versican in promoting [75] or inhibiting [76] cancer development is based on its vast network of interactions with other ECM molecules and cell-surface proteins including HA, multiple types of cell adhesion receptors and molecules, growth factors and their receptors and chemokines [74,81]. Versican is expressed as five isoforms (V0-V4) [77], V0 and V1 being the most prominent in cancer tissues as reviewed by Ricciardelli and colleagues [82]. Isoforms V0 and V1 have been indicated e.g. in promoting glioma [83] and melanoma cell behaviour [84] towards more malignant phenotype via their capability to interact with TGF–2 and the hyaluronal cell surface receptor CD44, respectively. Also other modular ECM PGs such as

**Table 1: Extracellular matrix PGs and HA in tumour microenvironment.**

<table>
<thead>
<tr>
<th><strong>SLRPs</strong></th>
<th><strong>Molecule</strong></th>
<th><strong>Core protein size (kDa)</strong></th>
<th><strong>GAG type (number of GAGs)</strong></th>
</tr>
</thead>
<tbody>
<tr>
<td>Decorin</td>
<td>40</td>
<td>CS/DS (1)</td>
<td></td>
</tr>
<tr>
<td>Biglycan</td>
<td>40</td>
<td>CS/DS (1-2)</td>
<td></td>
</tr>
<tr>
<td>Asporin</td>
<td>43</td>
<td>none</td>
<td></td>
</tr>
<tr>
<td>Fibromodulin</td>
<td>58</td>
<td>KS (4)</td>
<td></td>
</tr>
<tr>
<td>Lumican</td>
<td>38</td>
<td>KS (2-3)</td>
<td></td>
</tr>
<tr>
<td><strong>Modular PGs</strong></td>
<td><strong>Versican</strong> (isoforms V0-V4)</td>
<td>370, 262, 180, 72, 115 (V0-V4)</td>
<td>CS/DS (0-23)</td>
</tr>
<tr>
<td>Aggrecan</td>
<td>210-250</td>
<td>CS (100) KS (30)</td>
<td></td>
</tr>
<tr>
<td>Brevican</td>
<td>50,80,145</td>
<td>CS (0-5)</td>
<td></td>
</tr>
<tr>
<td>Perlecan</td>
<td>400-467</td>
<td>HS (3-10)(CS, DS, hybrid HS/CS, CS/DS chains or GAG-free glycoprotein)</td>
<td></td>
</tr>
<tr>
<td><strong>HA</strong></td>
<td><strong>Very low, medium and high HA species</strong></td>
<td><strong>Molecular weight (kDa)</strong></td>
<td><strong>Repeating glucuronid acid and N-acetylglucosamine disaccharides</strong></td>
</tr>
<tr>
<td></td>
<td>50, 300, 800, 2000-3000</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

*HA is the only GAG synthesized in a free form lacking a protein core.*
aggreca[n 85,86] and brevican [87] have been indicated with cancer development through their differential expression in cancer compared to healthy tissue. With aggrecan, it has been shown that in laryngeal squamous cell cancers the amount of aggregable aggrecan is excessively lost which is associated with cartilage destruction and linked with progression of the cancer [86].

Furthermore, chondrosarcoma cells are known to experience altered pattern of aggrecan mRNA splicing compared to normal cartilage. This is associated with transformation of chondrocytes into malignant cells [85]. Regarding brevican, its expression is upregulated in brain tumours including astrocytoma where it represents one of the ECM molecules responsible for invasiveness of the malignant cells and thus provides potential target for treatment for cancer [87]. Furthermore, in glioma the enhanced cell adhesion and migration of cancer cells have been shown to be dependent on proteolytic cleavage of brevican via activating e.g. EGF signaling resulting in secretion and accumulation of fibronectin [88]. Concerning the HSPG perlecan, its expression has been shown to be involved directly in the tumour progression through its ability to store and capture growth factors such as basic fibroblast growth factor (bFGF) [32,73]. In addition, in various cancer types including melanomas, the expression of perlecan has been shown to mark for aggressive phenotype [73].

Hyaluronan, a versatile non-sulfated GAG, which is produced by hyaluronan synthases 1-3 (HAS-1-3), consists of repeating glucuronic acid and N-acetylgalactosamine disaccharides. It can be found abundantly in most tissues of the human body [89,90]. HA does not have a core protein but it can bind to a variety of PGs namely hyalectans via the N-terminal domains of these PGs and act as an organizer of pericellular and extracellular matrices [31,90]. There is a vast amount of evidence ascertaining the role of HA in several cancer types such as prostate [91], ovarian [92], breast [93] and thyroid cancers [94] with stromal accumulation typical for progressed and poorly differentiated tumors indicating poor prognosis.

Specifically, breast and lung cancers exhibit more pronounced HA expression in the invasion front than in central tumour areas [95,96], and the metastatic score of breast cancer can even be assessed on the basis of HA metabolism [97]. The expression of HA has typically been linked with tumour progression via various mechanisms such as increased motility, invasive phenotype, proliferation and stimulation of growth factor production emphasizing a central role for HA as a modifier of tumour cell behaviour [98,99]. Binding of HA to its receptors, hyaladherins, e.g. CD44 and different isoforms of RHAMM [receptor for hyaluronan-mediated motility] [100,101] activates intracellular signaling leading to bidirectional information flow between the ECM and the cell genome in a phenomenon called dynamic reciprocity [102,103]. More precisely, CD44/HA binding leads to e.g. release of Basic Fibroblast Growth Factor (bFGF) and TGF-1 [99] whereas RHAMM/HA interaction activates Focal Adhesion Kinase (FAK) and extracellular signal-regulated kinase (ERK1/2) [104]. More examples concerning the possible roles of HA in desmoplastic reaction and tumourigenesis could also be introduced.

In addition to HA, also other GAGs, namely chondroitin, dermatan, and heparan sulfates which normally exist as side chains of PGs have been shown to variously accumulate in the microenvironment of a number of tumours including breast and ovarian cancers and thereby influence tumourigenesis [105-107]. However, it is noteworthy that the accumulation of GAGs does not necessarily promote tumourigenesis but can prevent it as has been shown for heparin-like polysaccharides [106].

Desmoplastic Reaction and Development of New Oncological Pharmacotherapies in the Future

As brought up above, drastic changes in the composition of ECM macromolecules including specific species of ECM PGs and also HA can take place in tumour microenvironment. These changes can lead to the formation of fibrotic response called desmoplastic reaction which can have various effects on tumourigenesis. The fibrotic response can also drastically restrict the use of pharmacotherapies, i.e. if the drug does not reach cancer cells, it cannot kill them despite of its efficacy in cell culture conditions [108]. Fibrosis in desmoplastic reaction causes problems to the delivery of anticancer drugs particularly in pancreatic [10,16] and liver cancers [15].

For example, in pancreatic cancer, the fibrotic tissue surrounding the tumour mass can account for >80% of the tumour volume [109]. In that perspective, inhibition of the synthesis of accumulating ECM macromolecules or their degradation could assist drug availability to the target tissue. Antifibrotic therapy on pre-neoplastic diseases such as liver cirrhosis [110] and pancreatic fibrosis [111] has already been promisingly tested in animals, but their real benefits in clinical trials wait to be evaluated. Furthermore, in malignantities like pancreatic ductal carcinoma and liver cirrhosis, the desmoplastic reaction also contains vast amounts of HA resulting in exceptionally high interstitial fluid pressure in solid tumour tissue, which in turn can prevent perfusion and diffusion of small molecule therapeutics [112]. Therefore, degradation of HA or restriction of its synthesis could be a rationale way to improve drug availability in these diseases. This kind of approach has already been successfully applied in the treatment of diabetes where the use of hyaluronidase can accelerate insulin exposure [113]. Also the use of 4-methylumbelliferone, a HA synthesis inhibitor has been shown to reduce HA induced fibrosis and to possess also other antitumour effects in hepatocellular carcinoma [114]. As it is natural, additional examples regarding oncological therapies that focus on ECM of tumours could be presented [115].

Conclusion

As discussed above, cancer cells create their own microenvironment. Desmoplastic reaction is a phenomenon where a dense fibrotic tissue rich in collagen and other ECM macromolecules including specific species of matrix PGs and HA around the tumour mass is observed. This reaction can either restrict or promote tumour progression. Desmoplastic reaction has even been suggested to precede tumourigenesis. Although much is already known about the possible mechanisms how individual ECM macromolecules involved in desmoplasia influence tumour cell behaviour, the importance of desmoplastic reaction as a whole in tumourigenesis still remains to be resolved. Nevertheless, targeting individual ECM macromolecules involved in desmoplastic reaction offers new promising ways to influence tumour progression via modulating e.g. growth factor activity. In the future novel drugs targeted to the microenvironment of cancers represent a direction of anticancer therapy that cannot be overlooked, especially in highly lethal cancer types such as pancreatic cancer associated with harmful desmoplasia. Furthermore, gaining better understanding of the process behind desmoplastic reaction and its influence on cancer progression would improve the development of even more effective cancer treatments.

References


