Hyaluronan-CD44 Interactions in Cancer: Paradoxes and Possibilities

Bryan P. Toole

Abstract

Hyaluronan is a prominent component of the micro-environment in most malignant tumors and can be prognostic for tumor progression. Extensive experimental evidence in animal models implicates hyaluronan interactions in tumor growth and metastasis, but it is also evident that a balance of synthesis and turnover by hyaluronidases is critical. CD44, a major hyaluronan receptor, is commonly but not uniformly associated with malignancy, and is frequently used as a marker for cancer stem cells in human carcinomas. Multivalent interactions of hyaluronan with CD44 collaborate in driving numerous tumor-promoting signaling pathways and transporter activities. It is widely accepted that hyaluronan-CD44 interactions are crucial in both malignancy and resistance to therapy, but major challenges for future research in the field are the mechanism of activation of hyaluronan-CD44 signaling in cancer cells, the relative importance of variant forms of CD44 and other hyaluronan receptors, e.g., Rhamm, in different tumor contexts, and the role of stromal versus tumor cell production and turnover of hyaluronan. Despite these caveats, it is clear that hyaluronan-CD44 interactions are an important target for translation into the clinic. Among the approaches that show promise are antibodies and vaccines to specific variants of CD44 that are uniquely expressed at critical stages of progression of a particular cancer, hyaluronidase-mediated reduction of barriers to drug access, and small hyaluronan oligosaccharides that attenuate constitutive hyaluronan-receptor signaling and enhance chemosensitivity. In addition, hyaluronan is being used to tag drugs and delivery vehicles for targeting of anticancer agents to CD44-expressing tumor cells. (Clin Cancer Res 2009;15(24):7462–8)

Background

The importance of the micro-environment in tumor progression is now well established (1–4). Hyaluronan is a prominent component of this micro-environment in most malignant tumors, both in the pericellular milieu immediately surrounding tumor cells and in the tumor stroma; its association with either compartment can be prognostic for tumor progression (5, 6). A major receptor for hyaluronan, CD44, is currently prominent in the cancer literature because it is a common marker for “tumor-initiating cells-cancer stem cells” (CSC) in human carcinomas. Even though the nature of CSCs is highly controversial, there is a reasonable consensus that CD44-expressing subfractions of many human carcinomas are highly malignant and resistant to therapy, properties that are frequently associated with CSCs (7, 8). Surprisingly, however, the functions of CD44 and its hyaluronan ligand in the properties of these particular cells have received little attention in the literature. The functional dynamics of hyaluronan and its receptors, especially CD44, have recently been reviewed in detail with respect to cancer (6, 9–17). In this brief overview, I will summarize my view of the current state of our knowledge of the functions of hyaluronan-CD44 interactions in cancer and the mechanisms whereby these interactions influence a large number of signaling pathways and cellular behaviors. Some of these activities are summarized in Fig. 1.

Hyaluronan. Hyaluronan (also hyaluronic acid or hyaluronate) is a very large, linear, negatively charged polysaccharide, which is composed of repeating disaccharides of glucurionate and N-acetylglicosamine. Hyaluronan is produced by three hyaluronan synthases (Has1/Has2/Has3), which are integral plasma membrane proteins whose active sites are located at the intracellular face of the membrane (18). Newly synthesized hyaluronan is extruded as it is elongated, and then targeted to the cell surface or to pericellular and extracellular matrices. Hyaluronan is distributed ubiquitously in vertebrate tissues but is especially concentrated in regions of cell division and invasion (19). In adult tissues such as synovial fluid, cartilage, and dermis, it clearly plays a structural role that depends on its unique hydrodynamic properties and its interactions with other extracellular matrix components. On the other hand, hyaluronan has an instructive, cell-signaling role during dynamic cell processes such as morphogenesis, inflammation, wound repair, and cancer, wherein hyaluronan-receptor interactions are activated and collaborate in driving numerous signaling pathways (11, 20, 21). In addition to signal transduction, hyaluronan-receptor
interactions participate in at least two other important physiological processes, viz., endocytosis of hyaluronan and assembly of pericellular matrices (19, 22, 23).

**Hyaluronan receptors.** Hyaluronan interacts with several cell surface receptors, including CD44, Rhamm, LYVE-1, HARE/stabilin-2, and Toll-like receptors-2 and 4 (20, 24, 25). CD44 is widely distributed but particularly important in the immune system and inflammatory processes (24, 26), as well as in diseases such as atherosclerosis and cancer (21, 27). Unlike CD44, LYVE-1, and HARE, Rhamm does not belong to the "link module" family of hyaluronan-binding proteins. Rhamm can be present either in the cytoplasm or on the cell surface, and is an important factor in cell motility in wound healing and cancer (10). LYVE-1 is a close relative of CD44, which is mainly restricted to lymphatic vessel and lymph node endothelia, but its function is not well established (25). HARE/stabilin-2 is a scavenging receptor that clears hyaluronan and other glycosaminoglycans from the circulation (28). The Toll-like receptors recognize hyaluronan fragments during inflammatory events (24).

The major receptors implicated in cancer are CD44 and Rhamm. I will focus on CD44 in this short review, but it is important to note that CD44 and Rhamm can exhibit both cooperative and interchangeable signaling functions. For example, interactions at the plasma membrane between CD44 and Rhamm have been shown to activate CD44 signaling through ERK1/2 and promote cancer cell motility (10). In some cases, e.g., in animal models of autoimmune diseases, Rhamm can compensate for CD44, a very important consideration when interpreting experiments in CD44-null mice (29). CD44 is a single chain, single-pass, transmembrane glycoprotein, which is very widely expressed in physiological and pathological systems. CD44 was first characterized through the confluence of several areas of investigation, including hyaluronan-cell interactions, lymphocyte homing, and cell adhesion (30), and its role in these phenomena is now well established (26,
Although CD44 arises from a single gene, numerous transcripts are formed by alternative splicing. "Standard" CD44 comprises of the constant, nonvariant exon products, whereas "variant" isoforms arise by splicing of numerous additional exon products into a single site within the membrane-proximal region of the ectodomain (31). Carcinoma cells typically produce several variant forms of CD44 as well as standard CD44, whereas some tumor types, e.g., gliomas, produce mainly the standard form (32). All forms of CD44 include an N-terminal, membrane-distal, hyaluronan-binding domain that has significant homology with the hyaluronan-binding region, i.e., link module, of several other proteins and proteoglycans. Hyaluronan is the most widely studied ligand for CD44, but other ligands are clearly important. The best characterized among these are osteopontin and factors such as FGF and selectin ligands that recognize carbohydrate side chains covalently bound to CD44 (31, 33). One of the most puzzling aspects of CD44 physiology is "activation" with respect to hyaluronan binding and consequent signaling. Possible factors contributing to activation include post-translational modifications of CD44 such as glycosylation, CD44-cytoskeletal interactions, localization of CD44 within specialized domains in the plasma membrane, and the mode of pericellular organization and presentation of hyaluronan, but none of these is well established. Nevertheless it is clear that numerous cytokines, growth factors, and alterations in cell context can induce the events that result in activation (31).3

**Hyaluronan-CD44 signaling.** In several types of cancer cells, binding of hyaluronan to CD44 results in direct or indirect interaction of CD44 with signaling receptors, such as ErbB2, epidermal growth factor receptor (EGFR), and transforming growth factor-β (TGF-β) receptor type 1, and influences the activity of these receptors (15, 21, 34). It can also lead to interaction with and altered activity of nonreceptor kinases of the Src family or Ras family GTPases (11, 34). Complex formation with adaptor proteins such as Vav2, Grb2, and Gab-1 mediates interaction of CD44 with upstream effectors, e.g., RhoA, Rac1, and Ras, which drive intracellular signaling pathways (11, 34). Thus, hyaluronan-CD44 binding influences the activity of a variety of downstream signaling pathways, especially the MAP kinase and PI3 kinase-Akt pathways, and consequently promotes tumor cell proliferation, survival, motility, invasiveness, and chemoresistance (11, 15, 20, 21, 31, 34). In addition, binding of hyaluronan to CD44 stimulates multidrug and metabolic transporters that are important in therapy resistance (11, 35–39) and presentation of proteases that facilitate invasion (31, 40). The bulk of current evidence indicates that these interactions involve specific variants of CD44 but the particular variant almost certainly depends on the type of tumor cell and stage of malignant progression, and in some cases standard CD44 rather than variant CD44 is critically involved. The mechanisms of regulation of these various interactions in different tumor cell types and stages are not well understood, but the widespread deregulation of many normal pathways in cancer cells most likely includes anomalous involvement of hyaluronan-CD44 interactions that operate normally in other contexts, such as embryonic development (19) and inflammation (24). A related possibility is that deregulated splicing in cancer cells (41) gives rise to CD44 variants that promote oncogenic events such as inappropriate Ras signaling (42), or binding of osteopontin, stromal growth factors, or proteases (31).

In addition to its action as a co-receptor or co-activator of membrane-associated signaling molecules, CD44 can influence cellular events such as tumor cell proliferation and motility through cross-linking to the actin cytoskeleton via ankyrin or members of the ezrin-radixin-moiesin family (11, 31, 40). The tumor suppressor Merlin most likely acts by inhibiting hyaluronan-CD44 interaction as well as displacing ezrin-radixin-moiesin proteins from the cytoplasmic tail of CD44. Release of Merlin suppression may trigger activation of hyaluronan-CD44 binding, which in turn leads to the formation of signaling complexes discussed above (40). Another mechanism whereby hyaluronan-CD44 interactions may lead to intracellular signaling is via intracellular cleavage of CD44, translocation of the cytoplasmic product to the nucleus and activation of transcription (43).

Many studies indicate that CD44 is localized at least in part to lipid micro-domains with the properties of lipid rafts, and associates indirectly or directly therein with signaling proteins and transporters. Most of these studies also show that CD44 is recruited into these domains in response to ligand interactions (11, 35, 44, 45). Moreover, endocytosis of hyaluronan and CD44 occurs from these domains (46). Given the large number of pathways affected by CD44, the possibility that indirect and direct interactions with a wide variety of effectors occur within such domains and that these domains are induced and/or stabilized by multivalent interactions of hyaluronan with CD44 provides a compelling postulate to guide current and future investigations, at least from this author’s perspective (see Fig. 1).

A puzzling aspect of many studies of hyaluronan-induced oncogenic signaling is their basis in experiments in which exogenous hyaluronan is added to cultured tumor cells. Although these studies have resulted in apparently solid data indicative of strong effects on the pathways in question, they are difficult to reconcile with the long history of safe use of hyaluronan in numerous reconstructive or regenerative capacities in human patients. For example, hyaluronan is employed widely in eye and knee surgeries and in prevention of adhesions (47, 48). Hyaluronan-based hydrogels are also being developed for a variety of purposes including drug delivery, encapsulation of progenitor cells, and tissue engineering (49, 50). Such studies imply that the oncogenic effects of hyaluronan only occur in the context of the tumor micro-environment and that stromal hyaluronan, as well as tumor cell-produced hyaluronan, play an important role in tumorigenesis, a conclusion supported by correlative studies of numerous human tumor types (6). Strong supporting evidence for the tumor-promoting effects of hyaluronan comes from studies in which tumor hyaluronan levels and interactions with receptors were manipulated in *vitro*. These studies are discussed briefly below.
models of several tumor types. The approaches used include manipulation of levels of hyaluronan and perturbation of endogenous hyaluronan-receptor interactions by a number of methods (e.g., see Fig. 2; refs. 13, 21). However, it has become evident that turnover of hyaluronan by hyaluronidases is an essential aspect of the promotion of tumor progression by hyaluronan and that the balance of synthesis and degradation is critical (12, 16). Recent work, in which hyaluronan synthesis was up-regulated conditionally in mammary tumors that arise spontaneously in MMTV-Neu mice, highlights the importance of hyaluronan in tumor promotion, especially via recruitment of stromal cells and angiogenesis (13). Numerous studies have shown an important role for hyaluronan-CD44 interactions in recruitment or homing of various cell types, including circulating immune cells and precursor cells (26, 51). The MMTV-Neu studies (13) also confirmed the importance of hyaluronan in epithelial-mesenchymal transitions (EMT). A major defect in the Has2-null mouse is failure to undergo EMT during early cardiac development (52). Moreover, up-regulation of Has2 in phenotypically normal epithelium induces the characteristics of EMT, including anchorage-independent growth and invasiveness (53), two of the major properties of malignant cells.

Evidence for involvement of CD44 in tumor progression is also strong but very complex. Studies of tumorigenesis in CD44-null mice and manipulation of CD44 levels in various tumor systems have provided contradictory results, but treatments with CD44 antibodies and vaccines have shown the importance of CD44 in tumor growth and metastasis in mouse models of leukemias and carcinomas (14, 54–56). Many studies have implicated variants of CD44 rather than standard CD44 in tumor progression, but this depends on the stage of progression and type of tumor (14, 31). A striking development in recent years is the emergence of CD44 as a marker for subpopulations of several types of human carcinomas, often termed CSCs, which exhibit highly malignant and chemoresistant properties (7, 8). Interestingly, the characteristics of EMT have recently been linked to the properties of these cell subpopulations. A CD44+/CD24− subpopulation exhibiting CSC properties is induced by up-regulation of EMT-associated transcription factors in primary human breast epithelium, and a similar subpopulation with both EMT and CSC properties can be isolated from transformed epithelial cells (8, 57). Notably, these cells exhibit anchorage-independent growth of colonies in soft agar, a property that usually reflects resistance to apoptosis, which in turn is linked to chemoresistance. Numerous studies have shown that the CSC subpopulation of carcinomas and other tumor types is resistant to chemotherapeutic agents, most likely because of increased anti-apoptotic pathway activity and enrichment of multidrug transporters (8, 15, 57). Another important outcome of EMT is invasiveness (4, 58) and, accordingly,
CSCs have been linked to invasiveness and metastasis (7, 8, 59). As noted above, hyaluronan is closely associated with EMT, and these same properties of anchorage-independent growth, resistance to apoptosis, drug resistance and invasiveness are induced or increased by up-regulation of hyaluronan synthesis and reversed by antagonists of hyaluronan-CD44 interactions (15, 21). In particular, strong evidence has been published showing hyaluronan-dependent association of CD44 with receptor kinases (21, 31, 34) and transporters (15, 35–39), which are important in drug resistance and malignancy. Recently, we have examined hyaluronan-CD44 interactions in a CSC-like subpopulation of cells isolated from human patient ovarian carcinoma ascites. We found that the CSCs are enriched in receptor tyrosine kinases and ABC-family drug transporters, that these proteins are present in close association with CD44 in the plasma membrane of the CSCs, and that this association depends on constitutive hyaluronan interactions (60).

**Clinical-Translational Advances**

Although the published literature on hyaluronan-CD44 interactions in cancer is riddled with paradoxes and apparent contradictions, most investigators in the field agree that these interactions offer an important target for translation into the clinic. A frequently expressed concern is the widespread expression and functions of hyaluronan and CD44 in normal physiology. However, two observations provide promise that therapeutic interventions can be developed that target oncogenic events with some degree of specificity or differential sensitivity. First is the finding that many of the interactions described above involve variants of CD44 that are amplified greatly in many tumor types in comparison to normal processes (14, 32, 41). Second is the nature of activation of hyaluronan-CD44 interactions in malignant tumor cells. Although these processes may have overlapping features with immune and inflammatory pathways, there are also clear differences that may be possible to exploit. Some of the studies described above have used antagonists that may ultimately have therapeutic value (e.g., see Fig. 2), but these have not yet reached the clinic. Below I have summarized some approaches that seem to show promise in approaching this important objective.

**CD44 antibodies and vaccines.** Several studies have shown that administration of antibodies against CD44 inhibits tumor growth and progression. For example, injection of monoclonal antibodies against CD44 that block binding of hyaluronan inhibit invasion of mouse lymphoma cells into lymph nodes (14). Antibodies against CD44 have also been shown to block homing and promote differentiation of acute myeloid leukemic stem cells, and consequently to eliminate tumor-initiating cells (54). Prolonged survival also occurred in mice with leukemic stem cells expressing BCR-ABL after treatment with CD44 antibody (55). Recently, a CD44 variant vaccine was shown to reduce mouse mammary carcinoma tumor growth and metastases (56). It is recognized in this field that these approaches will be greatly improved by tailoring antibodies and vaccines to specific variants of CD44 that are uniquely expressed at critical stages of progression of the cancer in question (14, 31). However, phase I trials in breast and head and neck carcinoma patients with an antibody against CD44v6 have been discontinued owing to toxicity (61, 62).

**Hyaluronidases.** Although constitutive hyaluronidase may promote the pro-oncogenic functions of hyaluronan, overexpression or exogenous administration of large amounts of hyaluronidase is usually inhibitory (12, 16, 17). Hyaluronidase has been used in the clinic for many years as an adjunct to chemotherapy in which it was believed to improve access of drugs to cancer cells through effects on cell adhesion and matrix barriers (12, 63). Highly purified recombinant hyaluronidase (64) is currently in phase I trials for patients with advanced solid tumors. Interestingly, hyaluronidase was also shown to sensitize mouse mammary carcinoma cells to chemotherapeutic drugs when the cells are cultured as drug-resistant spheroids (65), a technique now known to enrich for CSCs. Although hyaluronidase may act in part by reducing barriers to drug diffusion, it may also act via its oligosaccharide products, which have been found to inhibit constitutive hyaluronan-CD44 signaling, resulting in decreased cell survival and chemoresistance (see below; ref. 15).

**Small hyaluronan oligosaccharides.** Small oligomers of hyaluronan suppress anti-apoptotic signaling pathways in tumor cells and inhibit the activity of transporters that enhance resistance to therapeutic agents (15). Initially, the use of these oligomers was based on previous findings that oligomers consisting of 3 to 9 disaccharides bind CD44 monovalently (66) and displace hyaluronan polymer from membrane-bound receptors (67), but recent work has shown that they also inhibit hyaluronan synthesis (38). Treatment of tumor cells with these oligomers causes disassembly of CD44-transporter and CD44-receptor tyrosine kinase complexes, internalization of the disassembled components, and attenuation of function (38, 39, 44). Treatment in vivo with small hyaluronan oligomers suppresses tumor growth and/or induces tumor regression in experiments using xenografts of various tumor types, viz., melanoma, carcinomas, glioma, osteosarcoma, and malignant peripheral nerve sheath tumors (15, 21, 38, 68, 69). Notably, one of these studies showed significant effects on metastasis (68). Also, significant effects on tumor growth and invasion were seen when CSC-like subpopulations obtained from a glioma cell line (69) or from human patient ovarian carcinoma ascites (60) were used. Moreover, we have shown that systemic administration of suboptimal doses of hyaluronan oligomers sensitizes highly resistant, malignant peripheral nerve sheath tumors to doxorubicin treatment in vivo (38). Although this approach might be expected to interfere with all activated hyaluronan-CD44 interactions, malignant tumors seem to be far more sensitive than normal physiological processes.

**Targeting drugs to tumor cell CD44.** In addition to targeting of hyaluronan-CD44 interactions themselves, these interactions are being exploited for delivery of chemotherapeutic drugs and other anticancer agents to tumor cells. Many investigators have shown increased efficacy in cell and animal tumor models by conjugating drugs to hyaluronan or CD44 antibody or by incorporating drugs or small inhibitory RNAs into vehicles such as liposomes, hydrogels, and nanoparticles that have been decorated with hyaluronan or antibodies against CD44 (70). Initial trials in human patients with drugs conjugated to CD44 antibody have shown some promise, although they are complicated by various toxicities (70). Clearly, targeting to relevant variants of CD44 is a crucial aspect of this approach. It has also been found, however, that the enormous hydrodynamic domain encompassed by hyaluronan can be used to entrap drugs, without the need for chemical conjugation, and deliver them to CD44-expressing tumors.
Increased safety and efficacy of irinotecan, when combined with hyaluronan using this approach, have been shown in a pilot trial with colorectal carcinoma patients.

Disclosure of Potential Conflicts of Interest

B.P. Toole, inventor on patent related to content of article.

Acknowledgments

The author acknowledges the excellent work done by numerous investigators in the field. Because of restricted space, I have referred mostly to more comprehensive reviews that provide further details on primary sources of information and the various paradoxes and controversies in the field. I thank Drs. Eva Turley and Cornelio Tolg for critical reading of the manuscript.

References


70. Platt VM, Szoka FC, Jr. Anticancer therapies: targeting macromolecules and nanocarriers to hyaluronan or CD44, a hyaluronan receptor. Mol Pharm 2008;5:474–86.


Hyaluronan-CD44 Interactions in Cancer: Paradoxes and Possibilities

Bryan P. Toole


Updated version
Access the most recent version of this article at:
http://clincancerres.aacrjournals.org/content/15/24/7462

Cited articles
This article cites 69 articles, 19 of which you can access for free at:
http://clincancerres.aacrjournals.org/content/15/24/7462.full#ref-list-1

Citing articles
This article has been cited by 20 HighWire-hosted articles. Access the articles at:
http://clincancerres.aacrjournals.org/content/15/24/7462.full#related-urls

E-mail alerts
Sign up to receive free email-alerts related to this article or journal.

Reprints and Subscriptions
To order reprints of this article or to subscribe to the journal, contact the AACR Publications Department at pubs@aacr.org.

Permissions
To request permission to re-use all or part of this article, contact the AACR Publications Department at permissions@aacr.org.