The immunological effect of hyaluronan in tumor angiogenesis

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The relationship between the immune system and angiogenesis has been described in several contexts, both in physiological and pathological conditions, as pregnancy and cancer. In fact, different types of immune cells, such as myeloid, macrophages and dendritic cells, are able to modulate tumor neovascularization. On the other hand, tumor microenvironment also includes extracellular matrix components like hyaluronan, which has a deregulated synthesis in different tumors. Hyaluronan is a glycosaminoglycan, normally present in the extracellular matrix of tissues in continuous remodeling (embryogenesis or wound healing processes) and acts as an important modulator of cell behavior by different mechanisms, including angiogenesis. In this review, we discuss hyaluronan as a modulator of tumor angiogenesis, focusing in intracellular signaling mediated by its receptors expressed on different immune cells. Recent observations suggest that the immune system is an important component in tumoral angiogenesis. Therefore, immune modulation could have an impact in anti-angiogenic therapy as a new therapeutic strategy, which in turn might improve effectiveness of treatment in cancer patients.

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ANGIOGENESIS AND ANGIOGENIC FACTORS

Angiogenesis is defined as the formation of new blood vessels from pre-existing vessels. The angiogenic process may occur by four different mechanisms: sprouting, intussusceptions, elongation/widening and potentially, incorporation of circulating endothelial precursor cells into vessel walls. This process is critical during embryonic and fetal development, but also occurs as a physiological pathway in adults during wound healing, skeletal growth, menstrual cycle and pregnancy.

Physiological angiogenesis is a sequence of cellular events comprising vascular initiation, formation, maturation, remodeling and regression, which are accurately controlled to supply tissue requirements. Nevertheless, angiogenesis has an important role in different diseases including cancer. The biochemical stimulation of angiogenesis is performed by several angiogenic factors, being the most important: vascular endothelial growth factor (VEGF) and fibroblast growth factor-2 (FGF-2). Other angiogenic factors include angiopoietins, matrix metalloproteinases (MMPs), cadherins and integrins.1

VEGF is a major player in angiogenesis: VEGF-A is the main member of a gene family that also includes VEGF-B, VEGF-C, VEGF-D and placenta growth factor. VEGF interact with multiple receptor tyrosine kinases including VEGF receptor-1 (VEGFR-1) and VEGF receptor-2 (VEGFR-2). Placenta growth factor and VEGF-B selectively bind to VEGFR-1, whereas VEGF-C and VEGF-D primarily interact with VEGFR-3.2

VEGF-A–VEGFR-2 interaction induces a tyrosine kinases signaling pathway that stimulates proliferation, migration and production of several factors in endothelial cells (ECs). In turn this interaction stimulates vessel permeability (through endothelial nitric oxide synthase and nitric oxide), proliferation/survival (FGF-2), migration (ICAMs/VCAMs/MMPs) and finally differentiation into mature blood vessels.3 The FGF family consists of 22 known proteins, being FGF-1 (acidic FGF) and FGF-2 (basic FGF) the main members. In general, FGFs stimulates cellular functions by binding to cell-surface FGF-receptors in association with heparin proteoglycans. FGF-1 stimulates the proliferation and differentiation of the cell necessary for building an arterial vessel, including ECs and smooth muscle cells. Besides, FGF-2 promotes ECs proliferation and the tube-like structure organization. Even more, FGF-1 and FGF-2 are important players in wound healing process.3

TUMOR ANGIOGENESIS

Both in physiological and pathological angiogenesis, a cascade of several signals and cellular functions drive the establishment of new blood vessels, responding to an increased requirement of oxygen and nutrients.
A hypoxic microenvironment typically develops in pathological conditions such as cancer, enhancing VEGF production, which in turn cooperates with oncogenic events that promote angiogenesis. The hypoxic process is driven largely by the transcriptional activity of hypoxia-inducible factors, inducing the expression of VEGF-A and other pro-angiogenic mediators. Tumor vessels display disorganized structure and abnormal function, as tumor-associated ECs acquire changes in morphology, physiology, cytogenetics, epigenetics and gene expression. The irregular perfusion impairs oxygen, nutrient and drug delivery to the tumor. Finally, excess of pro-angiogenic molecules stimulated by hypoxia leads to an additional disorganization driving to malignant tumor cell selection and dissemination.

The role of tumor microenvironment in angiogenesis

Tumor cells are able to modify the environment where they grow and these changes support the formation of a special microenvironment: ‘the tumor microenvironment’. Tumor cells interact with tumor microenvironment, constituted by cellular and non-cellular components (the extracellular matrix (ECM)). Besides, tumor cells induce a disorder of the microenvironment homeostasis leading to a sustained proliferative signaling, evasion of growth suppressors, resistance to cell death, invasion and metastasis, reprogramming energy metabolism, evading immune responses and inducing drug resistance and angiogenesis.

Non-cellular microenvironment: ECM

ECM contributes to the angiogenesis process by multiple mechanisms: (i) is able to regulate migration, invasion, proliferation and survival of ECs, (ii) provides a scaffold where mechanical guidance forces are established among distal ECs, providing an organized environment in the absence of cell–cell contact, (iii) serves as a control of lumen and tube formation and (iv) offers neovessel stability and maturation.

The ECM components that mainly interact with tumor cells are fibroblast, laminin, collagen, proteoglycans and glycosaminoglycans, such as hyaluronan (HA) (Table 1). HA is a crucial component of the ECM. Interestingly, it has relevant implications in physiological and pathological angiogenesis, performing an importing link between ECM–angiogenesis–immune cells–cancer.

Cellular microenvironment

The cellular tumor microenvironment is composed of (i) non-hematopoietic cells including ECs and tumor-associated fibroblasts and (ii) stem cells, circulating endothelial precursor cells and hematopoietic cells as well as immune cells, such as myeloid-derived suppressor cells (MDSC), tumor-associated macrophages, neutrophils, natural killer cells (NK), B- and T-lymphocytes, monocytes and dendritic cells.

Several lines of evidence indicate that different immune cells subsets contribute to blood vessel neoformation and remodeling. Even more, the immune cells functionally participate during tumor growth and progression (Table 2). These cells release pro-tumourogenic factors like cytokines and chemokines, ECM-degrading enzymes, reactive oxygen species and other bioactive molecules. Different immune cells share several angiogenic signaling pathways with other types of cells in tumor microenvironment, thus inhibition of one factor might lead to the compensatory upregulation of another angiogenic molecule release from the immune system, maintaining angiogenic activity and causing therapeutic failure in clinical treatment.

As immune cells are able to bind HA through different receptors, the interaction between HA, immune system and tumor angiogenesis will be discussed in the following paragraphs of this review.

HA–immune system–tumor angiogenesis

In normal tissues, HA synthesis and degradation are strictly balanced throughout HA syntheses and hyaluronidases, respectively. At homeostasis, HA high-molecular weight (HMW, ranging from \(0.5 \times 10^6\) to \(2 \times 10^6\) Da) is predominant, having hydrodynamic properties. Although the low-molecular weight (LMW, ranging from \(10^4\) to \(0.5 \times 10^6\) Da) form is present mainly during inflammation. It is well known that in malignant tumors the concentration of HA is usually altered in relation to normal tissues and in some tumors the HA level could be considered a predictor of malignancy. In fact, during cancer, HA is fragmented into LMW forms due to an imbalance between HA syntheses/hyaluronidases activity and by reactive oxygen species present in tumor tissues. Even more, HA LMW and fragments (< \(10^4\) Da) promote spreading by stimulating angiogenesis and creating a microvascular network in several tumors. Moreover, it is well documented that HA oligomers are angiogenic and enhance tumor invasiveness. On the other hand and despite of their angiogenic properties, exogenous administration of HA oligomers might inhibit tumor growth. Besides, HA HMW could promote a protective effect during tumor progress, delaying tumor growth in colon carcinoma and migration in fibrosarcoma.

Table 1 ECM components in cancer

| Collagens          | Glycoproteins (laminins and fibronectins) | Proteoglycans and Glycosaminoglycans |
|--------------------|------------------------------------------|--------------------------------------|
| Laminins are heterotrimeric glycoproteins that are composed of α, β and γ chains. They are primarily located in BMs and form networks with type IV collagen and nidogen. When laminins are cleaved by MMPs, their subunits stimulate cell migration and invasion. On the other hand, fibronectin is dimeric glycoproteins that are present in the ECM and in blood. They form fibrils and affect cell morphology, adhesion, migration and differentiation by binding to integrins. Binding domains on fibronectin were found to modulate the activity of VEGF and promote the interaction between integrin and VEGFR-2. Glycosaminoglycans (GAGs) are long, unbranched polysaccharides composed of repeating disaccharide units consisting of alternating uronic acids and amino sugars. Four classes of glycosaminoglycans have been identified: heparan sulfate, chondroitin sulfate/dermatan sulfate, keratan sulfate and HA. The main chondroitin-sulfate proteoglycan of noncartilaginous tissues is versican. Syndecans, glypicans and CD44 are cell-surface proteoglycans. All of them present relevance in cancer. Changes in expression of these molecules, as well as of enzymes involved in their biosynthesis and degradation, contribute to the different steps of tumor progression: proliferation, growth, invasion and metastasis. In therapeutics, targeting of GAGs and proteoglycans are highly promising. In the case of HA, it is an anionic, non-sulfated glycosaminoglycan, formed by alternating glucuronic acid and N-acetylglucosamine. One of the main components of the extracellular matrix, HA contributes significantly to cell proliferation and migration, and may also be involved in the progression of some malignant tumors. | The most abundant proteins in mammals ECM. Fibrillar collagens influence cellular functions through interactions with integrins, laminin and hepan-sulfate proteoglycans. Increased deposition of collagen is found during tumor angiogenesis. It is known that proteolytic fragments of type IV collagen can inhibit tumor angiogenesis, as well as a proteolytic fragment of the α1 chain of type XVII collagen. Four classes of glycosaminoglycans have been identified: heparan sulfate, chondroitin sulfate/dermatan sulfate, keratan sulfate and HA. The main chondroitin-sulfate proteoglycan of noncartilaginous tissues is versican. Syndecans, glypicans and CD44 are cell-surface proteoglycans. All of them present relevance in cancer. Changes in expression of these molecules, as well as of enzymes involved in their biosynthesis and degradation, contribute to the different steps of tumor progression: proliferation, growth, invasion and metastasis. In therapeutics, targeting of GAGs and proteoglycans are highly promising. In the case of HA, it is an anionic, non-sulfated glycosaminoglycan, formed by alternating glucuronic acid and N-acetylglucosamine. One of the main components of the extracellular matrix, HA contributes significantly to cell proliferation and migration, and may also be involved in the progression of some malignant tumors. |

Abbreviations: ECM, extracellular matrix; HA, hyaluronan; MMP, matrix metalloproteinase; VEGF, vascular endothelial growth factor.
several works have demonstrated that HA HMW is able to mediate cancer resistance in naked mole rats. This species of rodent present a high HA accumulation in skin, and the bases of its cancer resistance is because their fibroblast cells synthesize large concentrations of HA with a molecular mass higher (\(-1 \times 10^7\) Da) than other species.\(^{40}\) Thus, this controversial effects require further investigations in the HA science study.

During inflammation immune cells are able to bind HA, thus their response depend on the cell type, the HA molecular weight, HA-binding proteins and structures\(^{41}\) (Figure 1). For instance, an inflammatory microenvironment induces macrophages to bind HA possible to promote their migration, survival and proliferation\(^{41}\) and it was shown that LMW HA polarize macrophages to M1 phenotype \textit{in vitro}.\(^{42}\) On the other hand, HA present in tumor microenvironment serves as a signal for recruiting tumor-associated macrophages, which are key immune cells involved in tumor neovascularization related to HA.\(^{43}\) Tumor-associated macrophages suffer a metabolic adaptation in hypoxic areas and release pro-angiogenic factors: VEGF, FGF-2, PIGF, platelet-derived growth factor (PDGF).\(^{44-47}\) Also, these cells are able to downregulate the expression of angiogenesis inhibitors, such as vasohibin-2\(^{48}\) and stimulate the expression of angiogenesis-modulating enzymes, such as COX-2, iNOS and MMPs.\(^{47,49,50}\)

MDSCs have an important role in tumor angiogenesis, as they are able to incorporate into endothelial vessel, displaying an endothelial like morphology and expressing both VEGFR-2 and VE-cadherin, which are EC markers. Besides, MDSCs secrete angiogenic factors, such as MMPs, VEGF, basic FGF.\(^{23}\) There is no evidence about their capability to bind HA, although it has been demonstrated in liver that HMW HA-binding proteins and structures\(^{41}\) (Figure 1). For instance, an inflammatory microenvironment induces macrophages to bind HA, which leads to immune tolerance either through T-cell deletion or through the differentiation of regulatory or suppressor T cells. Mature (activated) antigen-loaded DC, can start the differentiation of antigen-specific T cells into effectors T cells. DCs capture tumor antigens released from tumor cells and cross-present them to T cells, contributing to tumor rejection.\(^{24}\)

### Natural killer cells (NK)

Provide rapid responses to transformed cancer cells. Infiltration of tumors with NK cells has been shown to represent a positive prognostic marker in colorectal,\(^{26}\) gastric\(^{27}\) and lung carcinoma.\(^{28}\)

### B cells

High numbers of B-lymphocytes have been found in aggregates with other immune cells at the inflammatory site in tumor tissues of various human cancers.\(^{29}\) The intratumoral presence of B cells has been correlated with enhanced survival in patients with ovarian\(^{30}\) and non-small lung cancer.\(^{31}\)

There are several subsets of T cells, within CD4\(^+\) (helper) and CD8\(^+\) (cytotoxic), that recognize antigens expressed in most tumor cells. Enhanced intratumoral CD8\(^+\) T-cell infiltration has been observed to be a positive prognostic marker in melanoma, head and neck, breast, bladder, ovarian, colorectal, prostatic and lung cancers.\(^{22}\)

### Macrophages

They adapt their phenotype to the dynamically changing microenvironment that they encounter. TAMs are derived from circulating monocytes or resident tissue macrophages, found within the stroma of many tumor types.

### Myeloid-derived suppress cells (MDSCs)

This cells are a heterogeneous type of immune cell, which have a defined immunosuppressive function. Different factors derived from tumor microenvironment allow their expansion, in turn MDSC inhibit immune attack affecting T-cell response by several mechanisms.\(^{23}\)

### Dendritic cells (DC)

Are essential to induce immunity against cancer. Immature (non-activated) DC can present self-antigens to T cells, which leads to immune tolerance either through T-cell deletion or through the differentiation of regulatory or suppressor T cells. Mature (activated) antigen-loaded DC, can start the differentiation of antigen-specific T cells into effectors T cells. DCs capture tumor antigens released from tumor cells and cross-present them to T cells, contributing to tumor rejection.\(^{24}\)

Abbreviation: TAM, tumor-associated macrophage.

### Table 2 Presence of immune cells in cancer

| Immune Cells | Function |
|--------------|----------|
| Natural Killer Cells (NK) | Provide rapid responses to transformed cancer cells. |
| B cells | High numbers of B-lymphocytes have been found in aggregates with other immune cells at the inflammatory site in tumor tissues of various human cancers. |
| Macrophages | They adapt their phenotype to the dynamically changing microenvironment that they encounter. |
| Myeloid-derived suppress cells (MDSCs) | This cells are a heterogeneous type of immune cell, which have a defined immunosuppressive function. |

### Notes

- HA fragments and oligosaccharides, but not HA HMW, induce maturation and activation of dendritic cell (DC) \textit{in vitro} and in tumor context.\(^{42,52-55}\) Even though DCs prime responses to tumor antigens, functions of these cells are altered by the tumor microenvironment. Immature DC can increase the expression of VEGF and interleukin-8 (IL-8) on hypoxic conditions,\(^{46}\) promoting a pro-angiogenic function in cancer.\(^{57,58}\) DCs in tumor environment are able to release pro-angiogenic cytokines: tumor necrosis factor-\(\alpha\), IL-8, osteopontin,\(^{57,59}\) VEGF, oncostatin M,\(^{60,61}\) controlling the process of DCs transdifferentiation to EC-like phenotype.\(^{51}\) Besides, tumor-derived factors such as hepatocyte growth factor,\(^{62}\) transforming growth factor-\(\beta\), prostaglandin E\(_2\),\(^{54}\) lactate\(^{65}\) and osteopontin\(^{66}\) modulate DCs maturation and thus their pro-angiogenic properties.

- There is no direct evidence that links HA with NK cells, but it was shown that LMW HA in combination with IL-2, IL-12 or IL-18 is able to enhance interferon-\(\gamma\) production.\(^{67}\) Separately, it is well documented that NK cells contribute to physiological vascular remodeling during the secretory phase of menstrual cycle and pregnancy, through secretion of cytokines and angiogenic growth factors.\(^{68}\) The role of NK in tumor angiogenesis has not yet been fully investigated. Bruno et al. have shown that in patients with lung cancer, tumor infiltrating NK cells exhibited an angiogenic phenotype associated with a production of VEGF, PI GF and IL-8.\(^{69}\) NK cells have a high cytolytic capacity against transformed cancer cells, playing an important role in tumor rejection, even though NK cells function is provably affected in tumor microenvironment.

- Naïve B cells bind HA in a lower degree in comparison to activated B cells. However, it was shown that HA could induce their activation.\(^{70}\) Angiogenic signals between tumor cells, ECs and B cells are mediated by STAT3 through the overexpression of multiple pro-angiogenic molecules, such as VEGF, hypoxia-inducible factor-1, FGF-2, MMP-9, MMP-2.\(^{47}\) Furthermore, activated STAT3 B cells are found in the surroundings of tumor vasculature, contributing to tumor development and angiogenesis. Therefore, STAT3 in B cells might work as potential therapeutic target for anti-angiogenesis therapy.\(^{71}\) In parallel, it was demonstrated that STAT3 activation through fibroblast growth factor receptor (FGFR) induces HA synthesis, contributing to tumor growth.\(^{72}\)

- T cells activation induces HA binding probably to guide them to the inflammation site.\(^{41}\) Even more, it was demonstrated that HA stimulates CD4\(^+\) CD25\(^+\) T cells to increase IL-2 production. An IL-2 sustain production is able to (i) extend T-cell proliferation and/or (ii) stimulate Treg cells to suppress the immune response.\(^{41}\) In parallel, a hypoxic tumor microenvironment recruits Treg cells which promote

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tumor angiogenesis through VEGF-A production and diminish T-cell function. Alternatively, T cells are able to recognize class I and II major histocompatibility complex-peptide and other co-stimulatory surface molecules express on ECs. Furthermore, T cells are able to interact with ECs through cytokines and cell–cell contact, modulating blood vessel formation and remodeling, blood flow, fluidity and ‘permselectivity.’ T cells can synthesize pro-angiogenic factors, such as FGF-2 and heparin binding epidermal-like growth factor. Alternatively, T-cell cytokines, tumor necrosis factor-α, transforming growth factor-β and INF-γ were reported to have anti-angiogenesis properties in tumors such as brain and ovarian carcinoma.

The exact mechanisms by which HA can contribute with tumor angiogenesis has not yet been well elucidated. It is possible to suggest one mechanism linked to the immune system. As it was mentioned, HA oligomers and LMW are able to induce tumor angiogenesis, as well as inhibiting tumor growth and inducing active immune responses. To understand these apparent contradictions, further investigations will be necessary regarding the signals triggered by HA and its cellular receptors in tumor context.

**Modulation of intracellular signals by CD44**

CD44, the major receptor of HA, includes a family of plasma membrane glycoproteins, which are involved in several cellular functions, such as cell–cell and cell–matrix adhesions, cell migration, proliferation and lymphocyte homing. CD44 is expressed in a variety of cells including tumor cells. Elevated CD44 expression was correlated with poor prognosis in many cancers, such as lung, ovarian, breast, colorectal, gastrointestinal neuroendocrine tumor. Even more, CD44 is emerging as a metastatic tumor marker. As it has been stated HA–CD44 interactions have an important role in: (i) development, (ii) inflammation, (iii) T-cell recruitment and activation and (iv) tumor growth and metastasis. CD44 has a cytoplasmatic domain with phosphorylation sites (Ser291, Ser325 and Ser316) that are regulated when ligands bind to transduce signaling and select its downstream effectors. These interactions and modulations generate multiple cellular functions such as cytoskeleton activation, tumor cell adhesion, growth migration and invasion, leading to tumor progression and angiogenesis (Figure 2).

**HA–CD44 and RhoGTPases**

HA–CD44 induces activation of RhoGTPases signaling, which produce specific structural changes: as actin assembly, cytoskeleton reorganization, transcriptional activation, tumor cell growth, survival, migration and invasion. CD44 is linked to small GTP-binding proteins such as RhoA and Rac1. It has been shown that the interaction HMW-HA–CD44 leads to the activation of Rac1 initiating cortical actin formation and strengthening the connection between ECs. On the other hand, LMW HA binds to CD44 and triggers a series of events that promote the formation of RoA-GTP complex, which stimulates the serine/threonine kinase, leading to actin stress fiber formation and disruption of the EC barrier. These events triggered by LMW HA cause the disruption of the endothelium guiding to an important event in tumor angiogenesis and cancer metastasis. In immune cells, GTPases (RhoA, Rac1 and Rac2) have important roles in migration, adherence, chemotaxis and phagocytosis/endocytosis. However, it is not yet elucidated how the HA–CD44 interaction modulates RhoGTPases signaling in immune cells and its relation with the synthesis and function angiogenic factors.

**HA–CD44 and PI3K**

HA induces the formation of a complex that contains CD44, ErbB2, phosphatidylinositol 3-kinase (PI3K), ezrin...
and the chaperone protein HASP90. It was found that this complex is assembled in lipid rafts and it enables signaling through ErbB2 promoting cell survival through PI3K activation. This pathway has a central role in oncogenesis by the regulation of intracellular processes: cell survival, apoptosis, cell growth, angiogenesis, motility and chemoresistance. HA is able to modulate the PI3K/AKT pathway in different cancer types, such as lymphoma, colon and lung carcinoma. Activation of the PI3K/AKT pathway in tumor cells can increase VEGF secretion by hypoxia-inducible factor-1 and modulate the expression of other angiogenic factors: nitric oxide and angiopoietins. There is strong evidence that suggests that the PI3K/AKT pathway modulates angiogenesis though it is not well elucidated yet how HA is involved in this angiogenic pathway.

In inflammatory context PI3K/AKT controls several functions, for example hematopoietic stem cells survival, and T- and B-cell development. There is no evidence that connects this pathway in immune cells with CD44 and HA, but it is possible to hypothesize that HA would modulate this signaling by CD44 present in immune cells.

HA–CD44 and β-catenin. Another interesting outcome in the HA–CD44 signaling is the involvement of β-catenin, the main protein of the canonical Wnt pathway. The Wnt signaling pathway controls cell polarity, self renewal and proliferation. In absence of Wnt ligands, β-catenin is phosphorylated at N-terminal-specific residues by a destruction complex conformed by GSK3 kinase and the adenomatous polyposis coli protein among other complexes and it is sent to ubiquitination and degradation. Otherwise, when Wnt ligands bind to Frizzled receptors and low-density receptor-related protein (LRP) co-receptors, β-catenin remains unphosphorylated at GSK3 phosphorylation sites, which prevents its destruction, then it translocates to the nucleus and activates the transcription of target genes through the interaction with ‘lymphocyte enhancer factor’/‘T-cell factor’ nuclear transcription factors. β-Catenin-T-cell factor/lymphocyte enhancer factor complexes are involved in the transcription of cyclin D1 and c-myc target genes among others, which are important regulators of cell cycle and proliferation. In addition, β-catenin takes part in adherent junctions through binding to E-cadherin. Phosphorylation at specific C-terminal residues decreases β-catenin affinity for E-cadherin and β-catenin facilitates signaling transduction.

Preliminary data of our laboratory shows Wnt/β-catenin pathway activation and β-catenin relocalization in murine models of prolactinomas. Third South American Symposium in Signal Transduction and Molecular Medicine. Argentina. 2015. Interestingly, β-catenin signaling is a critical event in ErbB2-mediated mammary tumor and other tumor progression. Moreover, ErbB2 tyrosine kinase, modulated by HA–CD44 interaction, promotes β-catenin phosphorylation, which destabilizes E-cadherin-β-catenin complexes leading to a decrease of cell adhesion and transcriptional upregulation of T-cell factor/lymphocyte enhancer factor transcription, deregulating the cell cycle progression of ovarian tumor cells.

Furthermore, Wnt signaling regulates CD44 expression and thus its function, and also CD44 induces β-catenin instability reducing its nuclear accumulation. These reciprocal regulations between CD44 and Wnt/β-catenin result interesting in handling potential therapeutic targets, to inhibit metastasis, drug resistance and recurrent processes.

Several authors have investigated the involvement of Wnt/β-catenin signaling in vasculogenesis and angiogenesis. Franco et al. reviewed the emerging concepts of the role of Wnts, their receptors and signaling pathways in regulating differential behavior and/or cell function during vascular development. Indeed, Wnt pathway
regulates the expression of several angiogenic factors like VEGF-A\textsuperscript{97} and IL-8\textsuperscript{98} in different biological contexts, raising the possibility that it could promote both angiogenesis and sprouting of the pre-existing vessels.

It has been stated that treatment with IFN\textgamma, an inhibitor of the canonical Wnt signaling, as well as the knocking down of the Wnt pathway component BCL9 in different cancer models, decrease VEGF and CD44 protein and mRNA levels, suggesting a role of these factors regulated by Wnt in carcinogenesis.\textsuperscript{99-101} However, the relationship between the CD44-β-catenin crosstalk and angiogenesis is yet poorly understood.

Regarding to the roles of the Wnt/β-catenin signaling in immune cells, several functions have been described for this pathway, as the modulation of CD8\textsuperscript{+} T-cell differentiation and memory formation and regulation of CD4\textgamma T-cell polarization and survival.\textsuperscript{102} In the case of DCs, β-catenin in a Wnt-dependent and -independent manner modulates immunity and tolerance.\textsuperscript{103} Nevertheless, it is yet to be determined the mechanisms triggered by HA–CD44–β-catenin in immune cells.

**Modulation of ECM-degrading enzymes**

It has been reported that the interaction between HA and CD44 can induce, in tumor cells, the expression and activation of several ECM-degrading enzymes, such as MMP-2, MMP-7, MMP-9, EMMPRIN, cathepsin and MT1-MMP. This regulation has an important role in tumor cell behavior allowing matrix degradation, invasion and metastasis and also promoting angiogenesis during tumor progression.\textsuperscript{104-108} MMPs are important contributors to angiogenesis allowing new capillary formation and VEGF release from the ECM stores.\textsuperscript{109} The immune cells are able to produce different type of MMPs, in turn it is possible to suggest that HA may modulate MMPs in these cells. Once MMP-9 is stimulated, it activates the latent form of transforming growth factor-β modulating ECs behavior and inflammatory responses.\textsuperscript{110}

**Modulation of intracellular signals by TLR4**

Toll-like receptors (TLRs) are transmembrane proteins that recognize not only highly conserved molecules present on microbes, known as ‘pathogen associated molecules patterns’, but also responds to self-molecules produced after cell damage or death, identified as ‘damage-associated molecular patterns’. Damage-associated molecular patterns include a variety of different molecules present in the host. For example, LMW HA functions as a pro-inflammatory damage-associated molecular pattern as it can modulate DC, stimulate the release of pro-inflammatory cytokines and promote proliferation. Several of these pro-inflammatory effects of LMW HA are due to the interaction with TLR4.

It has been reported that the interaction of LMW HA with TLR4 induce specific intracellular signaling pathways in immune cells and ECs.\textsuperscript{54} TLR4 activation by LMW HA has a critical role during DC maturation, inducing the signal transduction pathway involving p38/p42-44 MAPKs and NF-kB.\textsuperscript{54} Besides, macrophages and tumor cells treated with LMW HA resulted in the activation of NF-kB, a major component of TLR4 signaling pathway, and lead to the transcription of MMPs (MMP-2) and cytokines (IL-8, IL-12 and tumor necrosis factor).\textsuperscript{54,111-113} This promotes motility and invasiveness in a TLR4 dependent manner.\textsuperscript{113} Even more, it was reported that LMW-HA fragments were found to interact with TLR4 in human dermal microvascular ECs inducing the release of IL-8.\textsuperscript{114} In parallel, the activation of NF-kB was found to stimulate angiogenesis by inducing expression of IL-8 and VEGF.\textsuperscript{115}

Despite there is not enough evidence that HA–TLR4 has a role in modulating angiogenesis, this mechanism cannot be ruled out. Our preliminary results show that HA acts as a modulator of VEGF expression in spleen cells (Spinelli et al. Hyaluronan as a modulator of VEGF expression in immune cells. 10th International Conference on Hyaluronan, Italy, 2015). In parallel, we are investigating the role of HA and its receptors, CD44 and TLR4, in the regulation of tumor angiogenesis by immune cells.

**Modulation of intracellular signals by RHAMM**

The receptor for HA-mediated motility (RHAMM, CD168) is a cytoplasmatic protein that is unconventionally exported to the cell surface.\textsuperscript{116} Intracellular RHAMM is associated with the centrosome and mitotic spindle.\textsuperscript{117} Extracellular RHAMM is a HA-binding protein that links and activates CD44 resulting in stimulation of the Ras/Ras1,2 pathway,\textsuperscript{118} a cascade that functions in cellular proliferation and is often deregulated in cancer. It was shown that in normal lymphocytes, RHAMM is mostly intracellular and surface RHAMM is almost always absent.\textsuperscript{119} On the other hand, RHAMM is expressed both as an intracellular and as a cell-surface protein in B lineage cancers, with important functional roles\textsuperscript{119}; motility and malignant spread,\textsuperscript{120} and also modulation of apoptotic and cell cycle progression pathways.\textsuperscript{116}

There are several studies that show that RHAMM is expressed in EC and it participates in modulating angiogenesis in EC. For example, it was shown that the RHAMM receptor is overexpressed in neovessels and it might be responsible for enhancing EC activation via HA oligosaccharides.\textsuperscript{121} RHAMM is necessary for EC migration\textsuperscript{122} and downregulation of RHAMM exerts a negative effect on HA-induced EC tube formation.\textsuperscript{123} Even more, it was shown that RHAMM stimulated by HA activates Rac1,\textsuperscript{124} a protein involved in the hypoxia-induced production of VEGF and VEGF-R.\textsuperscript{2,125,126}

In an inflammatory context, RHAMM–HA interaction was found to regulate macrophage chemotaxis and motility in lung injury.\textsuperscript{126}

**ANGIOGENESIS, IMMUNE SYSTEM AND HA AS THERAPEUTIC TARGETS IN CANCER**

Inhibition of angiogenesis is an important strategy for cancer treatment. Extensive research has positioned VEGF, FGF, PDGF/ PDGFR-β and angiopoietins as therapeutic targets. As VEGF has a central role in promoting an angiogenic phenotype in most tumors, development of VEGF antagonists has become one of the central focuses for anti-angiogenic therapies. The first evidence that anti-VEGF might be useful in cancer therapy came from a mouse tumor model using a neutralizing antibody, that could prevent tumor growth in neuroblastoma in mice.\textsuperscript{127} After that, antagonists of VEGF and its receptors have been developed and evaluated for their ability to suppress tumor growth. The strategies involved are (i) anti-hypoxia-inducible factor reagents, (ii) VEGF antisense oligonucleotides, (iii) VEGF ribozymes, (iv) soluble VEGF receptors, (v) expression of a dominant negative mutant of VEGF receptors in target ECs, (vi) anti-VEGF receptor antibodies, (vii) VEGF receptor-mediated downstream signaling pathways and (viii) VEGF DNA vaccines.\textsuperscript{128-130} Several anti-angiogenic agents have been approved for clinical use: bevacizumab (monoclonal antibody that targets VEGF), sunitinib, sorafenib and axitinib (small molecules tyrosine kinase inhibitors including VEGF-R and FGF-R). However, these drugs have not demonstrated the expected effectiveness in cancer treatment. Expression patterns of pro-angiogenic factors can be altered along tumor progression and single angiogenic factor antagonists may
encounter drug-resistance problems or activation of compensatory pathways.

Recent studies suggest that anti-angiogenic therapies are efficient to reverse some levels of immunosuppression in patients with cancer; therefore the link between anti-angiogenic strategies and immune system might lead to the appearance of new therapies, for example metastatic renal cancer patients treated with bevacizumab and interferon-α presented a superior progression-free survival than those treated with interferon-α monotherapy.\(^1\)

Furthermore, different anti-cancer therapies that are being consider are those that target other aspects related to HA. HA production, for example, through chemical inhibitors of HA synthesis like 4-Methylumbelliferone, an orally dietary supplement. It would be interesting and valuable to evaluate HA-related strategies in the context of anti-angiogenic therapy and immunotherapy.

**CONCLUSION AND FUTURE DIRECTIONS**

In summary, immune cells could modulate angiogenesis in different tumor contexts, having a crucial participation in neovascularization, beyond the known mechanism that involves EC. Thus, the factors and mechanisms that allow immune cells to acquire pro-angiogenic functions will be an important tool to improve and develop anti-angiogenic therapies in cancer. It is possible to hypothesize that abnormal ECM, is in part responsible for this unsuccessful response of the immune system. We also considered that HA, as a component of ECM, is able to modulate angiogenesis and immune responses, working as a linker in tumor microenvironment.

**CONFICT OF INTEREST**

The authors declare no conflict of interest.

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