Galectin-1 in Pancreatic Ductal Adenocarcinoma

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pancreatic ductal adenocarcinoma Galectin-1 tumor microenvironment angiogenesis

immune modulation

1. Galectin-1: Structural Complexity, Biological Function, and Emerging Role in Cancer Treatment

Galectin-1 is a unique and critical member of the galectin family, characterized by its ability to bind to βgalactoside-containing glycoconjugates ^[1]. Existing as a homodimer with 14-kDa subunits, it can be found in various tissues and demonstrates different subcellular localizations, such as in the cell nucleus, cytoplasm, cell surface, and extracellular matrix ^{[2][3]}. Gal-1 plays significant roles in normal and pathological conditions, being involved in diverse cellular processes such as cell cycle control, migration, adhesion, immune responses, apoptosis, and inflammation ^[4]. Its complex interactions can be sugar-dependent or sugar-independent and may be inhibited by specific compounds like thiodigalactoside (TDG) or lactose ^[4].

The molecular structure of Gal-1, encoded by the LGALS1 gene at 22q13.1, features a complex β -sandwich composition with two antiparallel β -sheets formed by five (F1-F5) and six (S1-S6a/b) strands, respectively ^[5]. In humans, it exists as a dimer stabilized by hydrophobic interactions within its core, formed by key residues from both subunits ^[1]. Additionally, a well-defined hydrogen bond network is established among specific residues ^[6]. Notably, the presence of six cysteine residues in the Gal-1 sequence makes it susceptible to oxidation, which might lead to changes in its physiological activity and function ^[7]. Refer to **Figure 1** for a visual representation of the Gal-1 molecular structure and its various cellular localizations, underscoring its diverse and crucial roles in cellular mechanisms.



Figure 1. Three-dimensional representation and subcellular localization of Gal-1 homodimer. (**A**) A detailed 3D model of the Gal-1 homodimer, accentuating its β -sandwich configuration. This structure comprises two antiparallel β -sheets, the first containing five strands and the second with six. The intricate architecture of Gal-1 is key to its binding properties and biological functions. (**B**) Schematic illustration of Gal-1's diverse cellular localizations, with depictions in the nucleus, cytoplasm, cell surface, and extracellular matrix. These varied locations emphasize Gal-1's multifaceted roles in regulating vital cellular processes including cell cycle control, migration, adhesion, immune responses, apoptosis, and inflammation.

Galectin-1 in Oncology: From Tumor Progression to Therapeutic Potential

Galectin-1 consistently manifests heightened expression within tumor tissues, encompassing not only the neoplastic cells but also the accompanying stromal cells ^[8]. Notably, it plays a significant role in the carcinogenesis of colorectal tumors ^{[9][10][11]}. Such heightened levels are often indicative of a more aggressive tumor behavior, promoting the development of a metastatic phenotype ^[12]. Beyond its association with tumor malignancy, Gal-1 exerts a significant influence in fostering tumor angiogenesis through modulation of endothelial cell activity, and it enhances the ability of tumor cells to evade immune surveillance ^{[5][13]}. Intriguingly, beyond its role in malignancies, Gal-1 has a contributory role in other inflammatory conditions of the gastrointestinal tract and in diseases that have recently gained significant attention, such as COVID-19 and ulcerative colitis ^{[14][15]}.

Gal-1's prominence in a variety of human cancers underscores its pivotal role in oncology ^[12]. Its diverse interactions in cancer, from nurturing metastatic tendencies to promoting tumor angiogenesis and helping tumors evade the immune system, accentuate its integral position in the complex landscape of cancer biology ^{[16][17]}. Given its profound influence, targeting Gal-1 might emerge as a promising strategy for therapies across multiple cancer forms ^[18].

2. Galectin-1 Expression in Normal and Pathological Pancreatic Tissues

In normal pancreatic tissues, Gal-1 expression manifests in a nuanced manner ^[19]. While most regions of the normal pancreas display weak or absent Gal-1 staining, in situ hybridization has identified pronounced Gal-1 mRNA signals in pancreatic nerves ^{[20][21]}. Additionally, immunohistochemical analyses have detected subtle Gal-1 immunostaining in select fibroblasts located within the intra- and interlobular stroma of pancreas ^[20]. The heightened expression of Gal-1 in both nerves and fibroblasts may suggest their potential roles in intercellular communication or tissue repair processes, pointing to a specialized function of Gal-1 within these specific cell types in the pancreas.

In pathological contexts of the pancreas, there is a marked upregulation of Gal-1 expression ^{[22][23]}. For instance, the studies of Wang et al. discerned moderate to intense Gal-1 expression predominantly in the fibroblasts of chronic pancreatitis specimens ^[23]. Furthermore, Pancreatic Stellate Cells/fibroblasts, pivotal in mediating desmoplastic responses across a spectrum of pancreatic diseases—from chronic pancreatitis (CP) and intraductal papillary mucinous neoplasms (IPMN) to pancreatic intraepithelial neoplasia (PanIN) and pancreatic ductal adenocarcinoma (PDAC)—exhibit pronounced Gal-1 expression ^[24]. Intriguingly, while ductal cells remain devoid of Gal-1, its presence is concentrated within the cytoplasm, nuclei, and extracellular matrix of Pancreatic Stellate Cells ^[24]. This elevated expression in Pancreatic Stellate Cells underscores Gal-1's potential role in modulating stromal responses, intercellular interactions, and possibly the progression of pancreatic pathologies.

2.1. Galectin-1 Expression in Pancreatic Ductal Adenocarcinoma

The role of Gal-1 in PDAC has garnered significant attention in the research community, shedding light on its specific localization and implications in the pathology of the disease ^[5]. In PDAC, a distinct concentration of Gal-1 is observed within stromal myofibroblasts, especially within activated PSCs embedded in the tumor-associated stroma ^[20]. The expression of Gal-1 in these stromal constituents suggests its potential involvement in stromal cell activation, which may promote tumor desmoplasia and progression ^[17].

Notably, poorly differentiated tumors exhibit an ascending gradient of stromal Gal-1 staining, indicative of its probable role in tumor aggressiveness or dedifferentiation ^[25]. Although initial research highlighted Gal-1's predominance in the tumor stroma, emerging studies have pinpointed its presence in cultured human pancreatic cancer cells ^{[22][26]}. A particular focus has been on the cell membrane localization of Gal-1 during cell migration ^[26]. Insights from specific mouse models have further identified robust Gal-1 staining in both tumor epithelia and stromal compartments ^[27]. Such findings hint at a possible reciprocal interaction between pancreatic tumor cells and their stromal environment. Specifically, the redistribution of Gal-1 to the cell membrane in epithelial cells upon stromal interaction could facilitate cell–cell communication or promote invasive tendencies ^[19].

Furthermore, the conspicuous expression of Gal-1 in the stromal milieu, potentially originating from its secretion by pancreatic cancer cells, might suggest a paracrine modulation of the desmoplastic reaction characteristic of PDAC

^[26]. Overall, the intricate expression patterns and localization dynamics of Gal-1 in PDAC highlight its significance in the disease's biology and the potential avenues for therapeutic interventions.

2.1.1. Galectin-1 mRNA Expression in Pancreatic Cancer Tissue

Analyses of Gal-1 mRNA levels within pancreatic cancer tissues have unveiled stark contrasts when juxtaposed with healthy controls ^[20]. Employing Northern blotting techniques, pancreatic cancer specimens exhibited markedly amplified Gal-1 mRNA levels ^[20]. Quantitative assessments discerned that the mRNA concentrations in these malignant samples were approximately 11.9-fold that of non-cancerous tissue, affirming the strong linkage between elevated Gal-1 mRNA expression and the intricacies of pancreatic cancer progression ^[20].

2.1.2. Galectin-1 Protein Expression in Pancreatic Cancer Tissue

Quantitative analysis reveals a substantial surge in Gal-1 protein concentrations in pancreatic cancer specimens, registering about 8.6-fold the levels seen in typical pancreatic tissues ^[20]. Detailed assessments identified pronounced Gal-1 immunoreactivity within the fibrous bands of tissue encapsulating the tumor, markedly surpassing the faint signals in regular pancreatic stroma ^[20]. It is intriguing that despite such profound stromal expression, pancreatic cancer cells largely remained devoid of Gal-1, echoing patterns observed in certain other cancers, including colorectal carcinomas ^[10].

2.1.3. Serum Levels of Galectin-1 in Pancreatic Ductal Adenocarcinoma

Galectin-1 has been identified as a small soluble molecule that can be secreted into the extracellular space via a non-canonical secretory pathway ^[1]. In the context of cancer, including pancreatic ductal adenocarcinoma, increased levels of Gal-1 have been detected in both plasma and serum, in conjunction with overexpression in tumor tissues ^[24].

In assessing PDAC patients, plasma concentrations of Gal-1 are markedly higher than those observed in healthy controls, as quantified using Enzyme-Linked Immunosorbent Assay (ELISA) ^[24]. With a delineated cut-off value set at 17.7 ng/mL, there is a harmonized balance between diagnostic sensitivity and specificity, championing the utility of Gal-1 as a potential standalone biomarker for PDAC identification ^[24]. When juxtaposed against the benchmark tumor marker Carbohydrate Antigen 19-9 (CA19-9), Gal-1 showcases comparable diagnostic sensitivity and specificity. Leveraging both markers in tandem—Gal-1 and CA19-9—has been proposed to diminish the incidence of false negatives in PDAC diagnosis, thus bolstering the precision of preliminary CA19-9 assessments ^[24].

3. Galectin-1 Expression and Clinical-Pathological Parameters in Pancreatic Cancer

In the context of pancreatic cancer, Gal-1 overexpression has displayed a complex relationship with various clinical-pathological parameters. Although no significant correlations were found between Gal-1 overexpression and general patient characteristics, histopathological tumor parameters, tumor stage, or postoperative survival, a

trend was identified ^[20]. Specifically, higher expression levels of Gal-1 were observed in more dedifferentiated cancer samples (G3), suggesting a potential relationship with tumor differentiation ^[20].

More notably, the expression of Gal-1 in the stromal region of PDAC tissues has been associated with multiple negative factors, serving as an indicator of unfavorable prognosis. These factors include larger tumor size, lymph node metastasis, perineural invasion, and poorer differentiation ^[22]. Elevated Gal-1 expression in the stromal compartment of PDAC has been further linked with perineural invasion and poor prognosis in patients, emphasizing the potential clinical relevance of Gal-1 as a biomarker in pancreatic cancer ^[25].

3.1. Galectin-1 and Lymph Node Metastasis in Pancreatic Cancer

Galectin-1 expression in pancreatic cancer demonstrates a significant association with the disease's progression and metastasis, particularly in lymph node metastases ^[28]. Compared to normal pancreatic tissues, Gal-1 expression has been found to be substantially higher in both primary malignant tumors and metastatic tissues ^[28]. This increase appears even more pronounced in advanced lymph node metastases, highlighting a potentially critical role for Gal-1 in pancreatic cancer's invasive nature ^{[25][28]}.

The mechanisms underlying this association seem multifaceted. The high expression of Gal-1 in stromal cells, including activated PSCs, is known to induce epithelial–mesenchymal transition (EMT) in cancer cells ^[25]. This induction facilitates the acquisition of a metastatic phenotype, further characterized by elevated Gal-1 levels in cancer cells themselves. As a result, these cells exhibit enhanced invasive and metastatic abilities, promoting tumor progression ^[28].

3.2. Galectin-1 Expression and Prognosis in Pancreatic Cancer

Galectin-1 has gained prominence as a significant prognostic factor in pancreatic cancer, with various facets of its expression shedding light on patient outcomes ^[25]. One critical observation is the link between survival rates and Gal-1 expression levels ^[20]. Specifically, patients who exhibit robust Gal-1 expression tend to have a shorter overall survival span—a median duration of 14.1 months—compared to the 24.8 months observed in those showcasing weak expression ^[29]. Delving deeper into the statistical implications, a multivariate survival assessment applying the Cox proportional hazard model pinpointed pronounced Gal-1 expression as an independent harbinger of unfavorable prognosis, manifesting a relative risk of 4.676 ^[29].

Furthermore, insights from quantitative proteomic profiling have underscored Gal-1's detrimental role concerning survival rates ^[29]. The intensity of Gal-1 staining, particularly in cancer-associated stromal cells, emerges as a pivotal parameter ^[25]. When considered alongside factors such as the tumor category and lymph node implications, it steadfastly holds its ground as an independent prognostic determinant ^[25].

Adding to this narrative, a comprehensive meta-analysis has echoed these findings ^[30]. It accentuates the strong association between elevated Gal-1 concentrations and a decline in overall survival for pancreatic cancer patients, suggesting a hazard ratio (HR) of 4.77 ^[30]. Another intriguing avenue of investigation revolves around Gal-1's

potential as a predictor of extended survival in pancreatic carcinoma cases, with preliminary results indicating a sensitivity rate of 64% and a specificity rate of 90% ^[31].

4. The Multifaceted Role of Galectin-1 in Pancreatic Ductal Adenocarcinoma (PDAC) Progression and Pathways

Galectin-1, a salient member of the galectin family, has been identified as a vital modulator of multiple signaling pathways contributing to the progression of pancreatic ductal adenocarcinoma ^{[19][20][22][26][32][33][34][35]}. This molecule's complex involvement ranges from its significant effects on signaling pathways like ERK and Hedgehog-Gli to intricate interactions with molecules such as RAS, p16INK4a, MAPK, EGFR-Pdx1, and TGF-β1/Smad2 ^{[19][22]} ^[33]. Recognizing the diverse roles of Galectin-1 could potentially unlock new therapeutic avenues to combat this aggressive cancer, especially given its impact on pancreatic cancer cell migration, fibrosis, and the epithelial–mesenchymal transition (EMT) ^[34]. For a concise overview of Galectin-1's multifaceted involvement in PDAC progression and pathways, please refer to **Table 1** provided below.

Role/Area of Influence	Details/Interactions	Implications for PDAC
Signaling Pathways	ERK, Hedgehog-Gli, RAS, p16INK4a, MAPK, EGFR-Pdx1, TGF-β1/Smad2	Critical in modulation, proliferation, and disease progression
Chemokine Production	Modulates MCP-1, CINC-1 mRNA expression	Enhances inflammation, promoting tumor growth and metastasis
Hh-Gli Pathway Interaction	Regulatory control over genes for migration, adhesion	Vital for PDAC progression and malignancy
RAS Pathway Affinity	Binding to H-Ras and K-Ras	Potential driver in pancreatic cancer progression
p16INK4a Interplay	Influences Gal-1 levels, impacts anoikis	Key player in PDAC cellular dynamics
EGFR-Pdx1 Axis Influence	EGFR-Pdx1 Axis Influence	Direct/indirect impact on PDAC progression
TGF-β1/Smad2 in PSCs	Promotion of feedback loop for fibrosis	Potential target for halting fibrosis in pancreatic cancer
PDAC Cell Migration	Modulation of SDF-1 secretion in PSCs	Possible therapeutic potential in targeting metastasis
EMT Role in PDAC	Inhibition reduces EMT markers, initiates via p38 MAPK pathway	Inhibition reduces EMT markers, initiates via p38 MAPK pathway

Table 1. The multifaceted role of Galectin-1 in PDAC progression and pathways.

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The Hedgehog-Gli (Hh-Gli) pathway is a pivotal signaling mechanism, particularly in the realm of pancreatic carcinogenesis, playing a crucial role in cell proliferation, differentiation, and tissue patterning ^[41]. Against this backdrop, in-depth investigations involving PANC-1 cells have spotlighted Gal-1's interaction with this critical pathway ^[19]. Gal-1 exerts regulatory control over genes essential for cell migration, adhesion, and malignant transformation, many of which intersect with the Hh-Gli pathway ^[19]. By modulating Gal-1 levels, researchers observed discernible effects on the Hh-Gli signaling components: diminished levels curtailed the pathway's activity, while elevated Gal-1 levels amplified Gli transcriptional activities ^[19]. These insights emphasize the instrumental role of Gal-1 in modulating the Hh-Gli pathway and underscore its profound implications for PDAC progression.

The RAS pathway stands out in the progression of PDAC ^[42]. Within this framework, Gal-1 has demonstrated a substantial binding affinity to H-Ras and, to a lesser extent, K-Ras, suggesting its potential as a pivotal figure in the progression of pancreatic cancer by modulating the Ras-signaling pathway ^[43].

The tumor suppressor p16INK4a plays a pivotal role in the regulation of Gal-1 expression, unveiling a crucial interplay within the cellular milieu ^{[44][45][46]}. When p16INK4a is present, there is a marked increase in Gal-1 levels, coinciding with a heightened expression of the α 5 β 51 integrin on the cell surface. This, in turn, instigates significant alterations in cell glycosylation patterns ^{[44][45][46]}. One of the most consequential outcomes of this interaction is its impact on anoikis, a specialized form of programmed cell death. Anoikis acts as a safeguard mechanism, preventing detached cells from colonizing elsewhere and forming harmful growths ^[47]. The modulation of Gal-1 levels by p16INK4a, along with the subsequent changes in cellular behaviors, plays a pivotal role in the progression of pancreatic ductal adenocarcinoma.

The EGFR-Pdx1 axis is increasingly recognized as a critical component in the molecular landscape of PDAC ^[19]. The Epidermal Growth Factor Receptor (EGFR) and Pdx1 transcription factor both have pivotal roles in pancreatic cell differentiation and proliferation ^{[48][49]}. Their dysregulation can drive the initiation and progression of PDAC,

making the EGFR-Pdx1 axis a vital point of investigation for understanding the disease's pathogenesis ^[50]. Within this intricate network, Gal-1 stands out as a consequential modulator. Observations have consistently shown that when Gal-1 expression is diminished, there's a notable decrease in both EGFR and Pdx1 RNA levels in tumors ^[19]. This suggests a potential regulatory relationship, with Gal-1 serving as an upstream influencer on the EGFR-Pdx1 axis. By modulating the expression or activity of these key proteins, Gal-1 may directly or indirectly impact the trajectory of PDAC.

Lastly, Galectin-1's interaction with the TGF-β1/Smad2 pathway in PSCs promotes a feedback loop likely expediting fibrosis ^[32]. Targeting Gal-1 could, thus, serve as a novel strategy for halting fibrosis in pancreatic cancer, emphasizing its critical influence on disease progression.

4.1. Galectin-1's Influence on Pancreatic Cancer Cell Migration

Galectin-1's role in enhancing the migratory potential of pancreatic cancer cells, particularly through the modulation of stromal cell-derived factor-1 (SDF-1), has been delineated ^[34]. Within PSCs, the predominant fibroblastic population implicated in the fibrotic responses of PDAC, Gal-1 directly governs the secretion of SDF-1 ^[34]. The downregulation of Gal-1 in these PSCs notably attenuates PDAC cell metastasis, underscoring the therapeutic promise of targeting Gal-1. Intriguingly, this regulatory mechanism pivots on the activation of NF-κB within PSCs, which subsequently boosts SDF-1 synthesis ^[34].

SDF-1, with its primary expression localized to PSCs, plays a consequential part in pancreatic cancer advancement ^[51]. When SDF-1 engages with its receptor, CXCR4, predominantly present on pancreatic cancer cells, it modulates tumor cell proliferation, malignancy, and resistance to chemotherapy. Notably, the interruption of the SDF-1/CXCR4 signaling axis via CXCR4 antagonists diminishes the tumor cells' migratory and invasive prowess, underscoring the axis's centrality in PDAC progression ^[51].

4.2. Galectin-1 and EMT in Pancreatic Carcinoma

Galectin-1 holds a significant role in the metastatic evolution and disease trajectory of PDAC ^[12]. When Gal-1 was targeted for suppression using small interfering RNA (siRNA), a pronounced decline in its expression within pancreatic cancer cells ensued. This resulted in a tempered migration and invasion potential of these cells ^[28]. A notable shift in cellular adhesion dynamics further accentuated Gal-1's critical influence on the motility of cancer cells ^[28].

Epithelial–mesenchymal transition (EMT) represents a key cellular program where epithelial cells, typically characterized by strong cell-to-cell adhesion and apical–basal polarity, undergo transformation ^[52]. They shed these epithelial traits and embrace mesenchymal features, which include an enhanced migratory capacity, invasiveness, and resistance to apoptosis ^[53]. In the context of cancer, EMT is particularly alarming as it endows tumor cells with these aggressive traits, facilitating metastasis ^[54].

Upon Gal-1 inhibition, there was a clear reduction in markers characteristic of EMT. Key transcription factors such as Snai1 and Twist, which orchestrate the EMT process, were subdued ^{[28][55]}. Concurrently, the expression of MMP-10, an enzyme instrumental to tissue remodeling and cancer invasion, was mitigated ^[28]. This offers compelling evidence of Gal-1's potential in initiating EMT, possibly in collaboration with MMP-10 and via the p38 MAPK pathway ^{[28][55]}. In an intriguing observation, the elevated expression of Gal-1, especially from PSCs, corresponded with increased levels of mesenchymal markers. Conversely, there was a reduction in epithelial markers ^[55]. This molecular rearrangement underscores the notion that PSC-derived Gal-1 actively promotes PDAC invasion by amplifying the EMT phenomenon.

When PANC-1 cells were co-cultured with PSCs that overexpressed Gal-1, these cells underwent marked phenotypic alterations, suggesting a mesenchymal transition ^[55]. The enhanced invasive attributes of PANC-1 cells in this setup were further reflected by molecular markers, reinforcing the idea that PSC-sourced Gal-1 augments tumor invasiveness. Interestingly, this influence of Gal-1 on the EMT process might have its roots in the NF-κB signaling pathway, given the observed fluctuations in associated molecular markers when PANC-1 cells interacted with PSCs harboring variable Gal-1 concentrations ^[55].

5. Galectin-1 in the Tumor Microenvironment of Pancreatic Cancer

In pancreatic cancer, Gal-1 originates predominantly from tumor fibroblasts and stellate cells ^[20]. While these cells secrete Gal-1 that binds to the extracellular matrix (ECM), pancreatic tumor cells also express and secrete Gal-1, contributing to its stromal presence ^[56]. The paracrine uptake of this protein by endothelial cells further amplifies its reach and effects ^[19]. **Figure 2** visualizes this intricate interplay where Gal-1 orchestrates a series of events in the PDAC microenvironment, culminating in the profound desmoplastic reaction and subsequent ischemic conditions, thereby emphasizing the multifaceted role of Gal-1 in pancreatic cancer progression and microenvironment modulation.



Figure 2. Mutual reinforcement between Galectin-1 and Pancreatic Stellate Cells in PDAC microenvironment. The schematic illustrates the cascade initiated when PDAC (Pancreatic Ductal Adenocarcinoma) releases Galectin-1 (Gal-1), which activates and stimulates the proliferation of PSCs (Pancreatic Stellate Cells). These activated PSCs subsequently release a myriad of mediators, including IL-6 (Interleukin-6), IL-8 (Interleukin-8), IL-10 (Interleukin-10), and TGF-β (Transforming Growth Factor-beta). These mediators contribute to a series of downstream effects: Remodeling of the ECM (Extracellular Matrix), Collagen Synthesis, Conversion to CAF (Cancer-Associated Fibroblast) cells, and fostering ECM deposition, cumulatively leading to a pronounced desmoplastic reaction. This desmoplasia contributes to the ischemic environment characteristic of PDAC. Furthermore, the activated PSCs themselves release Gal-1, signifying a mutual reinforcement loop between Gal-1 and PSCs. The impact of Gal-1 is also discerned in PDAC cells, where it heightens the invasiveness, fortifying their migratory and invasive tendencies.

Gal-1 plays an instrumental role in the tumor microenvironment of PDAC ^[22]. Beyond stimulating chemokine production and accelerating PSC proliferation, it also amplifies the synthesis of key matrix components like collagen and fibronectin ^{[33][57]}. This drives a pronounced desmoplastic reaction, characterized by heightened stromal deposition and ECM remodeling, facilitating tumor growth and immune evasion ^[20]. When overexpressed in fibroblasts, Gal-1 not only accentuates the desmoplastic response but also fortifies the fibrotic barriers that hinder immune cell infiltration and compromise chemotherapy efficacy ^[56]. The depletion of Gal-1 in specific pancreatic tumors, in fact, results in reduced desmoplasia, underscoring its critical influence in shaping the tumor milieu ^[19].

Gal-1's Influence on Angiogenesis in PDAC

Pancreatic ductal adenocarcinoma displays a dense vasculature with poor perfusion and diminished integrity ^[58]. Despite its abundance, this vascular network fails to adequately deliver nutrients, oxygen, and drugs, largely due to the high interstitial fluid pressures stemming from fibrous interstitial fluid, which compresses blood vessels and hampers effective drug penetration ^{[59][60]}.

Galectin-1 plays a crucial role in fostering angiogenesis in PDAC, particularly through its interaction with the extracellular matrix ^[19]. This angiogenic promotion is intricately tied to its influence on tumor cell metabolism. Central to this influence is the "Warburg effect", wherein tumors preferentially undergo glycolysis even in oxygenrich environments. Through modulation of this metabolic shift, Gal-1 not only enhances nutrient supply via angiogenesis but also indirectly supports tumor immune evasion ^{[35][61][62]}.

The hypoxic tumor microenvironment, orchestrated by hypoxia-inducible factor-1 (HIF-1), stabilizes Gal-1 expression in conjunction with glucose transporter-1 (GLUT1) ^{[35][63]}. This dynamic fosters not only angiogenesis but also enhances tumor proliferation and metastatic potential. Within this context, Gal-1, similar to its counterpart galectin-3, propels glycolysis via pathways like the Phosphoinositide 3-kinase (PI3K) signaling cascade ^[35]. The resultant spike in lactate levels, emerging from glycolysis, acidifies the tumor environment ^[64]. This acidity not only promotes angiogenesis but also lures immunosuppressive components ^[64]. By enhancing the activity of enzymes linked to glycolysis, Gal-1 augments lactate production, facilitating the tumor's adaptation to hypoxic conditions, which in turn promotes angiogenesis and suppresses immune responses ^[35].

Empirical evidence solidifies Gal-1's significance in angiogenesis ^[43]. Enhanced Gal-1 expression in the tumor vasculature is associated with increased endothelial cell proliferation and migration, a relationship potentially modulated by H-Ras signaling ^[43]. Conversely, the diminished presence or absence of Gal-1 hampers these angiogenic processes ^[19]. Studies on Gal-1 deficient mice further support this, showcasing stunted tumor growth attributed to impaired angiogenesis ^[65]. In specific mouse models, lower Gal-1 levels were associated with reduced angiogenesis and decreased intraperitoneal hemorrhages, further underscoring Gal-1's essential role in angiogenesis and the broader spectrum of tumor progression ^[19].

6. Galectin-1 and Pancreatic Stellate Cells in PDAC Fibrosis

Pancreatic Stellate Cells are pivotal in engendering stromal fibrosis in PDAC ^[66]. They achieve this by remodeling the ECM and releasing a range of compounds, such as IL-1, IL-6, IL-8, IL-10, VEGF, Platelet-Derived Growth Factor (PDGF), and TGF- β , which play integral roles in collagen synthesis, and fostering ECM deposition ^{[32][67]}.

Galectin-1 is a characteristic product released by PSCs and plays a significant role in enhancing the function of these cells ^[68]. Laboratory investigations have shown that Gal-1 can boost the growth of PSCs and their production of collagen ^[33]. Interestingly, substances produced by PSCs also appear to stimulate the release of more Gal-1 ^[69]. This suggests a mutual reinforcement between Gal-1 and PSCs, where each enhances the activity of the other.

Within the context of PDAC, Gal-1 plays a crucial role in directing PSCs to produce cytokines like IL-6, IL-10, and IL-8 ^[35]. This action in turn speeds up the deposition of ECM, a process vital for tumor development. In animal studies, Gal-1 has been observed to enhance PSC-driven IL-10 production, resulting in a more fibrous tumor environment ^[68]. Additionally, under low-oxygen (hypoxic) conditions, Gal-1 further magnifies its influence by amplifying HIF expression, prompting PSCs to have an even greater fibrotic impact ^[70]. Studies in mice that lack Gal-1 have shown disruptions in the tumor stroma related to the Hedgehog (Hh) signaling pathway, emphasizing Gal-1's critical contribution to the fibrosis seen in PDAC ^[19].

Gal-1 is integral to the inflammatory responses that contribute to fibrosis in pancreatic cancer ^[71]. Notably, it catalyzes the activation of the NF-κB pathway, thereby intensifying inflammation. Such heightened inflammation propels fibrosis, which is primarily orchestrated by PSCs ^[33]. As these PSCs navigate the tumor microenvironment, they engage with immune cells, including macrophages and B lymphocytes. This cellular crosstalk magnifies the fibrotic landscape characteristic of PDAC ^[72]. In addition, Gal-1 disrupts the immune equilibrium between Th1 and Th2 cells in pancreatic malignancies. An outcome of this disruption could be a surge in IL-5, a cytokine implicated in the progression of tumor-associated fibrosis ^{[68][73]}.

Studies have highlighted a notable increase in Gal-1 expression in PSCs from PDAC samples when compared to normal pancreatic tissues ^[20]. When Gal-1's activity is deliberately curbed in human PSCs, these cells display diminished migration and invasive tendencies ^[74]. In contrast, introducing Gal-1 externally encourages cell growth and boosts collagen production in rat PSCs. These effects hint at the involvement of specific cellular signaling mechanisms, including the MEK1/2-ERK1/2 pathways, as well as the engagement of transcription factors like NF- κB ^{[33][57]}. With the accumulating insights, it becomes apparent that Gal-1, especially when expressed by activated PSCs, could serve as a potent target for immunotherapeutic strategies in treating pancreatic cancer.

6.1. Autocrine Effect of Gal1 on PSC Activation in PDA

Pancreatic stellate cells, when activated, play a significant role in the progression of pancreatic ductal adenocarcinoma ^[66]. As these PSCs become active, they undergo noticeable changes: they lose their fat droplets and show increased levels of α -smooth muscle actin (α -SMA), which is vital for the process of fibrogenesis ^[75]. Alongside these changes, there's a marked rise in Gal-1 expression, which transforms the appearance of these cells, making them resemble fibroblasts ^[22].

Gal-1, found in activated PSCs, plays a vital role in maintaining their active and fibrogenic characteristics ^[22]. Interestingly, when the levels of Gal-1 are reduced in human PSCs, these cells tend to become less active, almost returning to a dormant state ^[57]. This suggests that Gal-1 is crucial for keeping PSCs activated and supporting their transformation into fibroblast-like cells.

Gal-1 exerts a significant autocrine effect on PSCs, shaping their behavior and pivotal role in the progression of pancreatic cancer. Specifically, when Gal-1 binds to its ligands, it sets off a cascade of cellular signals ^[27]. A notable pathway influenced by this binding is the ERK pathway, crucial for fostering cell growth and ensuring

survival ^[69]. Another is the TGF- β 1/Smad signaling, which, when activated by Gal-1, propels PSCs into an enhanced state of activity ^[32]. In tandem, these signaling pathways dictate the transformation of PSCs, positioning them as central players in pancreatic cancer progression ^[69]. Delving into how Gal-1 modulates these pathways and the activation of PSCs provides a deeper understanding of the intricate interplay between PSCs and Gal-1 within the context of PDAC. Acquiring such insights is fundamental in demystifying the root causes and charting prospective treatment strategies for pancreatic cancer.

6.2. Paracrine Secretion of Gal-1 by PSCs in Pancreatic Cancer

Gal-1, secreted by activated PSCs, exerts powerful paracrine effects on surrounding cells, especially on surrounding pancreatic tumor cells and the immune milieu ^[69]. To put it simply, a paracrine effect means that a molecule produced by one cell can influence the behavior of nearby cells. Evidence from laboratory investigations has underscored that exposure to recombinant Gal-1 (rGal-1) accentuates the invasiveness of specific pancreatic cancer cell lines, such as CFPAC-1 ^[76]. Concurrently, media conditioned by human PSCs have been shown to bolster the migratory and invasive tendencies of the RWP1 pancreatic cancer cell lines—a phenomenon that attenuates significantly upon depletion of Gal-1 ^[27].

On a molecular echelon, the paracrine actions orchestrated by Gal-1 induce a cascade of cellular modifications within pancreatic carcinoma cells ^[27]. Notably, there's an upregulation of proteins like vimentin, coupled with a concomitant downregulation of E-cadherin, fostering an environment conducive for increased cellular migration and invasion ^[55]. Moreover, Gal-1, stemming from PSCs, amplifies the proliferative capacity of pancreatic tumor cells ^[25]. This is exemplified by enhanced cell multiplication when these tumor cells are co-cultured with PSCs exuberantly expressing Gal-1, in stark contrast to their counterparts with diminished Gal-1 levels ^[25].

6.3. Gal-1's Influence on PSC-Mediated MMP Activation in PDAC

In the intricate milieu of pancreatic malignancy, the role of Gal-1, predominantly secreted by PSCs, emerges as vital, especially in the orchestration of matrix dynamics ^[68]. Elevated Gal-1 expression in PSCs augments the secretion of matrix metalloproteinases (MMP-2 and MMP-9) ^[76]. These enzymes, known for their proteolytic functions, are instrumental in degrading the extracellular matrix (ECM) and cellular basement membrane (BM), thus facilitating tumor progression and invasion capabilities ^[77]. A notable observation was that of CFPAC-1 cells, which, when in co-culture with PSCs, showcased an increased MMP-2 and MMP-9 synthesis, coupled with accentuated proliferation and invasiveness ^[76]. This effect was pronounced when interacting with cancerassociated PSCs (CaPSCs) as opposed to their normal counterparts (NPSCs) ^[25]. However, this amplified activity seemed to diminish with the introduction of lactose, a recognized inhibitor of galectins ^{[25][76]}.

Delving further into the molecular intricacies, an engaging interplay between MMP-2, Tissue Inhibitor of Metalloproteinase-1 (TIMP-1), and Transforming Growth Factor-beta 1 (TGF- β 1) was discerned ^[32]. Gal-1's interaction with PSCs led to a discernible increase in MMP-2 levels and an even more substantial rise in TIMP-1 levels. This modulation suggested Gal-1's potential role in encouraging ECM protein synthesis, an indication of its

pro-fibrogenic tendencies ^[32]. To understand in greater detail, MMP-2 aids both in the proliferative activity and the degradation of type IV collagen present in the basement membrane ^[78]. Yet, the overarching expression of TIMP-1 serves as a counteracting force. This dynamic pushes the balance between ECM synthesis and degradation towards fibrogenesis, illuminating Gal-1's integral role in the genesis of pancreatic fibrosis ^[79].

6.4. Gal-1's Interplay with CAF in PDAC Development

In the intricate landscape of PDAC, cancer-associated fibroblasts (CAFs) play a quintessential role in fostering tumor fibrosis ^[80]. These CAFs, constituting a significant portion of the ECM, are primarily driven to activation under the influence of cytokines and agents such as TGF- β , IL-6, and IL-10 ^{[81][82]}. Upon activation, their physiological response leads to the secretion of matrix metalloproteinases (MMPs) ^[83]. This secretion acts as a catalyst for the degradation of the basement membrane, ensuing in comprehensive ECM remodeling. Concurrently, these fibroblasts potentiate fibrosis via the release of hyaluronic acid and collagen ^[84].

Delving into the relationship between Gal-1 and CAFs, it is evident that Gal-1 emerges as a significant pro-fibrotic mediator, meticulously orchestrating CAF activation. Particularly noteworthy is its ability to enhance IL-6 and IL-10 secretion in the PDAC milieu, thereby accelerating the activation of CAFs which in turn amplifies stromal tumor fibrosis. Insights from murine studies also spotlight Gal-1's potential in precipitating CAF activation via the Hedgehog signaling pathway ^[19]. Moreover, within the domain of PSCs, a surge in Gal-1 expression has been linked to an indirect activation of CAFs—this is predominantly mediated by an upregulated TGF- β expression, leading to a subsequent rise in MMP secretion and profound changes in the ECM ^[35].

7. Gal-1 and Immune Evasion in Pancreatic Ductal Adenocarcinoma (PDAC)

In the intricate realm of immunology, Gal-1 emerges as a molecule of paramount significance, endowed with robust immunoregulatory attributes ^[4]. It has been observed to exert a profound influence on neutrophil dynamics by curtailing their migration, thereby altering the innate immune response ^[56]. Furthermore, its role in macrophage plasticity is noteworthy; specifically, Gal-1 facilitates the transition of macrophages from the M1 phenotype, which is inherently pro-inflammatory, to the M2 phenotype, characterized by its immunosuppressive properties. This transformation, when contextualized, points toward Gal-1's overarching theme of tempering immune responses, as seen by its propensity to encourage the maturation of tolerogenic dendritic cells and to foster T-regulatory cell differentiation ^[35].

Delving into the context of pancreatic cancer, the role of Gal-1 becomes even more salient. Within the tumor microenvironment, it acts as a pivotal architect in establishing an immunosuppressive milieu ^[68]. One of the salient manifestations of this is seen in how Gal-1 deters T cell activation, tipping the scales towards an immunological state that is less reactive ^[27]. Gal-1 plays a noteworthy role in suppressing the immune response against tumors. Specifically, it acts by inducing apoptosis in T cells, which are vital components of the immune system responsible for recognizing and attacking tumor cells. By promoting the death of these T cells, Gal-1 effectively weakens the

immune system's ability to identify and combat tumor cells ^[85]. Beyond this direct impact on immune cells, Gal-1 also exerts influence on the broader immune environment. It actively modulates the secretion patterns of cytokines, which are signaling molecules that dictate immune responses. By altering these secretion patterns, Gal-1 creates an anti-inflammatory environment, favoring tumor survival and growth, and this modulation affects a wide range of immune cells, further emphasizing its significant role in cancer progression ^[68].

In the broader perspective of pancreatic pathologies, early indications of Gal-1 expression have been associated with stromal activation events such as inflammation and preneoplasia ^{[24][86]}. Its tangible presence in the stroma during ailments like pancreatitis or preceding malignant transformations signifies its role in creating an immune sanctuary ^[24]. This niche environment is conducive for immune evasion, laying a fertile ground for tumor inception and progression. Drawing a nexus between Gal-1-mediated stroma activation and the attainment of immune privilege paints a comprehensive picture of its centrality in pancreatic maladies, highlighting the molecule's potential as a therapeutic target. Following this intricate interplay, **Figure 3** provides a visual representation of the stages and processes where Gal-1 takes center stage, emphasizing its pivotal role in stromal activation, immune evasion, and the subsequent evolution of pancreatic cancer.



Figure 3. The immunosuppressive role of Gal-1 in PDAC. The illustration depicts the multifaceted impact of Gal-1 in mediating immune suppression and evasion within the PDAC microenvironment. Enhanced Gal-1 expression is shown to inhibit the chemotaxis and infiltration of neutrophils into PDAC tissue. The altered immune landscape is marked by a diminished T lymphocyte count, with a pronounced Th2 cytokine profile. Crucially, Pancreatic Stellate Cells (PSCs) expressing Gal-1 act as formidable barriers, obstructing T cell infiltration. Gal-1 further instigates

apoptosis in Th1 and Th17 cells, reinforcing its immunosuppressive role. The detrimental effects extend to the Natural Killer (NK) cells, where Gal-1 is seen to inhibit and inactivate them, largely due to the compromised IL-2 production and elevated IL-6 expression. The scene reveals an augmented presence of tolerogenic dendritic cells and MDSCs, underscoring the immune-evasive strategies employed. Finally, the PDAC is encapsulated by an abundance of tumor-associated macrophages (TAMs) of the M2 phenotype, with neighboring MDSCs, highlighting their synergistic role in thwarting T cell functionality and bolstering apoptosis. Note: PDAC (Pancreatic ductal adenocarcinoma), PSC (Pancreatic Stellate Cells), IL (Interleukin), Gal-1 (Galectin-1), CAF (Cancer-associated fibroblasts), MDSC (Myeloid-derived suppressor cells), M1 (Type 1 macrophage), M2 (Type 2 macrophage).

7.1. Galectin-1 Influence on T Cell Dynamics in PDAC

Acquired immunity plays a pivotal role in pancreatic pathologies, orchestrating complex interactions that influence disease progression and outcomes ^[87]. Galectin-1, a β -galactoside-binding protein, profoundly modulates T cell function in both physiological and pathological states ^[88]. Its influence is particularly pronounced in PDAC, where it emerges as a facilitator of tumor immune evasion ^[35]. Within the PDAC environment, Gal-1 takes center stage as a key agent promoting tumor immune evasion ^[35]. One of its striking features is its ability to curb T cell infiltration ^[56]. Specifically, while CD3+ T cells can be observed in the vicinity of malignant cells, they face a formidable barrier in penetrating the tumor parenchyma ^[62]. This suggests a significant role for Gal-1-expressing PSCs in shaping the tumor-immune landscape, ostensibly by impeding T cell access to the tumor core ^[56].

Delving deeper into the molecular intricacies of Gal-1, its adeptness in orchestrating T cell apoptosis is evident ^[29]. The molecule demonstrates multifaceted mechanisms in this regard. Its interaction with the CD45 molecule on T cells culminates in the degradation of fodrin, setting the stage for T cell phagocytosis and consequent apoptosis ^[89] ^[90]. Moreover, the binding of Gal-1 to receptors such as CD7 and Complement receptor 3 (CR3) augments this apoptotic cascade ^{[91][92]}. Notably, in the presence of Gal-1, quiescent T cells are propelled towards cell death via pathways such as the fas/caspase-8 axis, and intriguingly, Gal-1 can even induce the release of molecules like ceramide, which can activate other apoptosis circuits, including those involving caspase-9 and caspase-3 ^{[93][94]}. There is also an underpinning of the JNK/C-Jun/AP-1 pathway in Gal-1-mediated apoptosis ^[95].

Lastly, Gal-1's influence is not confined to direct cellular interactions but also extends to the modulation of the cytokine milieu in PDAC. The overexpression of Gal-1, particularly in PSCs, effects a paradigm shift in cytokine dynamics: it diminishes the expression of Th1 cytokines, such as IL-2 and INF- γ , while concurrently promoting Th2 cytokines like IL-4 and IL-5 ^[68]. This cytokine modulation, pivotal in shaping immune responses, can be partially reversed by attenuating Gal-1 levels ^[68]. A case in point is the observed suppression exerted by endogenous Gal-1 from PSCs on Th1 cytokines, a phenomenon particularly pronounced in human cancer-associated PSCs (hCaPSCs) ^[29]. Importantly, this shift from a Th1 to Th2 cytokine profile holds profound implications for tumor immunity, fostering an environment more amenable to tumor progression and potentially undermining anti-tumor immune responses.

7.2. Galectin-1 and NK Cell Dysregulation in PDAC

Natural Killer (NK) cells are innate immune effectors with the inherent ability to eliminate malignant, aged, and virus-infected cells through various cytokines and proteins ^[96]. However, within the hostile environment of PDAC, these formidable defenders become vulnerable due to the overexpression of Gal-1 ^[35].

Within the complex milieu of PDAC, Gal-1 plays an influential role, particularly concerning the modulation of NK cell activity ^[35]. Customarily, NK cell activation, which is facilitated by IL-2 and marked by the release of key entities such as Interferon- γ (IFN- γ) and tumor necrosis factor- α (TNF- α), is seen to be compromised in the presence of PDAC ^[97]. This anomaly is likely due to the ability of Gal-1 to curtail the production of IL-2, thus tipping the Th1/Th2 cytokine balance and diminishing NK cell activation ^[68]. Furthermore, Gal-1 seems to enhance IL-6 secretion, a factor which is known to negatively impact NK cell functionality ^[98].

Obesity, an often-discussed risk factor for PDAC, has been linked to elevated levels of Gal-1, resulting in increased adipocyte infiltration. This raised expression of Gal-1 coincides with heightened IL-6 levels, which in turn leads to a reduction in IFN-γ production, thus hindering NK cell operations ^[98]. Additionally, the presence of elevated Gal-1 also seems to promote the secretion of matrix metalloproteinase (MMP9) and indoleamine 2,3-dioxygenase (IDO), both of which are recognized for their inhibitory effects on NK cell efficiency ^[99]. The correlation between increased Gal-1 secretion and type-2 diabetes, commonly observed in PDAC patients, presents a significant concern ^[100]. This relationship intensifies insulin resistance, typical of type-2 diabetes, and boosts Gal-1 production, further dampening NK cell functionality in the face of PDAC ^[101].

Lastly, attention must be directed towards NK Group 2, Member D (NKG2D), a critical NK cell surface receptor. This receptor, vital for NK cell activation, interacts with MHC class I-related molecules ^[102]. Yet, in PDAC conditions, its expression appears reduced, a phenomenon potentially influenced by the hypoxic conditions amplified by Gal-1. In this scenario, Gal-1 is believed to foster a hypoxic environment, thereby augmenting Hypoxia-Inducible Factor (HIF) activity ^[103]. This escalation seems to inhibit NKG2D expression, consequently weakening the combative capabilities of NK cells against tumor formations.

7.3. Galectin-1's Role in TAM-Mediated Immune Evasion in PDAC

Tumor-associated macrophages (TAMs), specifically of the M2 phenotype, play a pivotal role in immune suppression within PDAC landscapes ^[104]. In synergy with myeloid-derived suppressor cells (MDSC), they inhibit T cell functionality and augment T cell apoptosis ^{[105][106]}.

Notably, a defining characteristic of PDAC, the activation of hypoxia-inducible factor-1 (HIF-1), shows a pronounced association with TAM expression ^[107]. This association is further marked by an enhanced lactate production, underscoring the intertwined nature of these phenomena ^[35]. During hypoxic events, the HIF-1 pathway is observed to escalate Gal-1 concentrations, an effect that seems to be further amplified by the concurrent H-ras pathway ^[108]. This surge in Gal-1 notably stimulates the production of vascular endothelial growth factor (VEGF) through the upregulated secretion of interleukin-6 (IL-6) ^[109]. In another significant sequence, upon lipopolysaccharide (LPS) exposure, Gal-1 is known to instigate ADAM10/17 production, which subsequently gives

rise to lactic acid ^[103]. This chain of events activates the nuclear factor erythroid 2-related factor 2 (Nrf2), influencing reactive oxygen species (ROS) generation and consequently steering TAM differentiation and augmenting VEGF secretion ^[109].

Moreover, an interesting interplay is evident when Gal-1 binds to Neuropilin1 (NRP-1) found on CAFs ^[110]. This association appears to facilitate VEGFR2 signaling, thereby dampening angiogenesis while enhancing the hypoxic attributes inherent to PDAC ^[111]. Such conditions are conducive for the activation of both TAM and MDSC, further strengthening the TAM-driven immunosuppression prevalent in PDAC. To encapsulate, Gal-1 appears to be a central figure in molding the immunosuppressive facet of PDAC, primarily through its influence on TAM activation and proliferation.

7.4. Gal-1's Influence on MDSCs in Pancreatic Ductal Adenocarcinoma

In the realm of Galectin-1 functionalities, its profound influence over immune cell infiltration in PDAC cannot be understated ^[4]. A cellular element under its considerable purview is the MDSCs, entities renowned for their pronounced immunosuppressive characteristics ^[112]. Within the context of Kras-driven PDAC, tumor cells exhibit a propensity to release granulocyte-macrophage colony-stimulating factor (GM-CSF), a molecule instrumental in choreographing the MDSC recruitment ^{[106][113]}. The pivotal orchestration of Gal-1 in MDSC accumulation is further illuminated by experimental observations wherein Gal-1-devoid mice depict a marked reduction in MDSC presence ^[27]. It behooves to comprehend the significance of MDSCs in PDAC, as their heightened presence often correlates with advanced tumor stages, underscoring the vital role they play in tumor progression and immune evasion ^[114].

7.5. Tolerogenic Dendritic Cell Modulation by Galectin-1 in PDAC

Galectin-1 emerges as a critical molecular entity in the landscape of PDAC, particularly in its influence on the differentiation of dendritic cells towards a tolerogenic disposition within the tumor microenvironment ^[115]. This pivotal differentiation underlines Gal-1's cardinal role in fostering an immunosuppressive milieu within PDAC, thereby bolstering tumor-driven immune tolerance ^[27]. Tolerogenic dendritic cells, characterized by their unique attributes, not only play sentinel roles in PDAC but also act as mediators, bridging the chasm between reactive and suppressive immune responses intrinsic to the tumor ecosystem ^[116]. The potential diminishment or absence of Gal-1 might perturb these finely tuned mechanisms, resulting in a conceivable augmentation in effector T cell populations within the tumor setting ^[27]. This potential shift underscores the profound implications of Gal-1's modulation on the immunological equilibrium of PDAC, highlighting its potential as a therapeutic target.

7.6. Gal-1's Modulation of Neutrophils in Pancreatic Tumors

Neutrophils, occupying a cardinal position within the tumor microenvironment of PDAC, are subjected to notable regulatory modulation by Gal-1 ^[117]. This molecule exhibits a salient propensity to impede the chemotaxis and consequent migratory abilities of these neutrophils ^[118]. Conspicuously, Myeloperoxidase (MPO) staining elucidates an enhanced neutrophilic presence in ductal malignancies concomitant with Gal-1 abrogation ^[19]. Such empirical observations underscore the imperative function of Gal-1 in safeguarding the immune privilege inherent

to pancreatic neoplasms, attenuating neutrophil-centric immunological activities, thereby accentuating its consequential role in the overarching therapeutic landscape of PDAC.

8. Gal-1 Inhibition in PDAC: Therapeutic Horizons

Gal-1, with its evident role in promoting immune evasion in pancreatic cancer, is being studied as a therapeutic target ^[35]. Depletion of Gal-1 has been linked with an increased influx of T cells and neutrophils in pancreatic tumors, shedding light on its potential to reinstate the body's immunological defenses against cancer ^[56]. Existing data indicate that anti-Gal-1 strategies, like inhibitors or monoclonal antibodies, exhibit anti-proliferative and anti-angiogenic effects across multiple tumor types, underscoring their potential in countering Gal-1-induced immunosuppression in pancreatic tumors ^[18][119].

Notably, the observed viability and fertility in Gal-1-knockout mice suggest a possible safety of treatments targeting Gal-1 ^[56]. This makes Gal-1 an interesting possible target for immune treatments against pancreatic cancer. But moving from lab studies to real-life treatments is not simple. Galectins, including Gal-1, are complex and need careful study before making specific drugs. Aspects like protein conformation, receptor glycan repertoire, cell type, and cellular conditions can alter the inhibitory effects of Gal-1. Therefore, using results from animal tests directly in humans may not always work. This is because humans and animals can have differences in how these proteins are expressed, their specific roles, and how the immune system works. Thus, it is important to be careful and study well before making new treatments ^[56].

However, the substantial expression of Gal-1 in human pancreatic cancer and its demonstrable influence on immune response enhancement, tumor cell proliferation reduction, and tumor stroma modification solidify its position as a prospective therapeutic candidate. To harness this potential fully, further research is imperative to refine the nuances of Gal-1 inhibition, enhance inhibitor specificity, and assess their effectiveness and safety in human trials. In this context, two avenues are emerging as particularly significant: the development of specific inhibitors and the promising intersection of Gal-1 with nano-oncology.

Inhibitors: lab trials have spotlighted a Galectin-1 inhibitor, LLS2, that augmented the efficacy of chemotherapy agent paclitaxel across multiple human cancer cell lines, encompassing pancreatic cancer cells ^[120].

Nanotechnology: Nano-oncology is carving out novel avenues to deploy galectin-based inhibitors or galectin ligand-augmented nanoparticles for diverse oncological applications. A recent study pivoted on Gal-1's upregulation in pancreatic cancer for the delivery of magnetic nanoparticles to cancer tissues ^[120]. These nanoparticles were adapted with glycosylated peptides from tissue plasminogen activator (tPA), a protein overexpressed in PDAC and recognized as a Gal-1 ligand ^[120]. Mouse PANC-1 xenograft magnetic resonance imaging elucidated a superior uptake of these tailored nanoparticles relative to non-specialized ones ^[120]. This endorses Gal-1's potential as a therapeutic vector in pancreatic cancer. Gal-1 directed nanoparticles might usher in groundbreaking strides in pancreatic cancer theranostics, merging diagnostic imaging with therapeutic intervention.

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