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REVIEW

The Role of STING-Mediated Activation of Dendritic Cells in Cancer Immunotherapy

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Abstract: The signaling pathway that comprises cyclic guanosine monophosphate–adenosine monophosphate (cGAMP or GMP-AMP) synthase (cGAS) and Stimulator of Interferon Genes (STING) is emerging as a druggable target for immunotherapy, with tumor-resident dendritic cells (DC) playing a critical role in mediating its effects. The STING receptor is part of the DNA-sensing cellular machinery, that can trigger the secretion of pro-inflammatory mediators, priming effector T cells and initiating specific antitumor responses. Yet, recent studies have highlighted the dual role of STING activation in the context of cancer: STING can either promote antitumor responses or enhance tumor progression. This dichotomy often depends on the cell type in which cGAS-STING signaling is induced and the activation mode, namely acute versus chronic. Of note, STING activation at the DC level appears to be particularly important for tumor eradication. This review outlines the contribution of DCs in cancer. We further highlight how the STING pathway plays an intricate role in modulating the function of DCs embedded in tumor tissue. Additionally, we discuss the strategies being employed to harness STING activation for cancer treatment, such as the development of synthetic agonists and nano-based delivery systems, spotlighting the current techniques used to prompt STING engagement specifically in DCs. **Keywords:** dendritic cell, cGAS-STING pathway, cancer immunotherapy, nanoparticle

Introduction

The paradigm of natural antitumor immunity involves the recognition, processing and presentation of cancer-associated antigens by dendritic cells (DCs) on major histocompatibility (MHC) complexes, triggering effective T cell-mediated antitumor immunity that results in killing of the cancer cells.^{1–4} Accordingly, DCs are highly specialized tissue-scanning cells equipped with a plethora of receptors to detect damage- or pathogen-associated molecular patterns (DAMPs or PAMPs). Although the immune system's task of filtering and responding to onco-signals emanating from cancerous tissue is an exceedingly complex one, the wide range of innate receptors expressed by DCs enables them to effectively recognize several by-products associated with tumor growth, such as tumor-derived DNA present in the tumor micro-environment (TME).^{5–7} The cGAS-STING pathway is an innate DNA-sensing mechanism which was originally described to mediate protective immune responses against infections but was recently found to play a crucial role in driving antitumor immunity. In particular, activation of STING signaling in DCs has the potential to generate spontaneous as well as long-lasting and highly specific antitumor immunity.^{8–10} This review focuses on the contribution of individual DC subsets to cancer defense and explores emerging developments in STING-mediated activation of DCs for novel cancer therapeutics.

The cGAS-STING Pathway and Its Effects in Cancer Cells

The cGAS-cGAMP-STING signaling pathway, discovered in 2013, is a promising therapeutic target to enhance antitumor immunity. Its activation induces an inflammatory response that converts "cold" tumors into "hot" tumors, ie, cancers that have higher infiltration of immune cells and consequently are more likely to exhibit higher response rates to currently used

immunotherapies such as immune checkpoint inhibitors (ICI).^{8,9,11} STING is a ubiquitous intracellular receptor that is located in the endoplasmic reticulum membrane and activated by binding of the agonist cGAMP. cGAMP is a naturally occurring cyclic dinucleotide (CDN) produced when the enzyme cGAS detects intracellular nucleic acids. Upon binding cGAMP, STING dimerizes and undergoes vesicular trafficking to the Golgi. This triggers downstream signaling which involves activation of TANK binding kinase 1 (TBK1), phosphorylation of interferon (IFN) regulatory factor 3 (IRF3), and induction of nuclear factor kappa B (NF-κB).^{12,13} As a result, type I IFNs (IFN I) are upregulated, leading to expression and secretion of pro-inflammatory factors that can affect the TME and influence disease progression.¹⁴⁻¹⁶ In cancer cells, the cGAS-STING pathway is frequently activated due to genomic instability caused by higher occurrence of mitotic errors compared to healthy cells. Consequently, tumors are rich in micronuclei, and chromatin bridges. Micronuclei consist of residual chromosomes or chromosome fragments that are surrounded by a nuclear envelope that persists after mitosis. These structures are prone to rupture, and by releasing nucleic acids into the cytosol, they have been reported to trigger STING activation.^{17,18} Interestingly, Flynn et al report that in spite of the ability of these structures to efficiently recruit cGAS, they fail to activate the cGAS-STING pathway.¹⁹ Instead, the authors convincingly show that chromatin bridges, single-handedly trigger cGAS activation, and subsequent IFN signaling.¹⁹ Similarly to micronuclei, these structures are also byproducts of mitotic processes which persist throughout interphase and can be strongly stretched, until they break due to tension caused by cell migration.¹⁷ Self-chromatin can induce STING activation, that in turn, has the potential to impede early tumor progression by upregulating inflammatory genes and by inducing apoptosis as well as cell cycle arrest, in a process mediated by IFN I and NF-KB.^{20,21} In melanoma, for example, restoration of STING signaling in tumor cells increases their immunogenicity and thus their recognition by T cells.²² Species-specific molecular determinants of self-DNA reactivity of cGAS were also uncovered using a melanoma model, which contributed to a better understanding of innate immunity involved in self-nonself discrimination. By engineering melanoma cells to express several species-specific cGAS transgenes with a doxycycline-inducible system, the authors showed that the key factor that protected mice from tumor-related mortality was the self-DNA reactive cGAS present in melanoma cells.²³ In fact, induction of STING in tumor cells can induce cell death, increasing the availability of tumor antigens to be recognized by antigen-presenting cells, such as DCs. Accordingly, besides the self-sustained protection mechanisms supported by STING activation in cancer cells, the continuous engagement of this receptor recruits various immune cells to the TME; therefore, enabling the cGAS-STING pathway in the immune compartment is of critical importance to overcome the generation of tumor-suppressive immune cells by the TME.^{24,25}

STING-Mediated Activation in DCs

Tumor-resident DCs are attractive targets for cancer immunotherapy because potent activation of DCs can initiate effective antitumor immune responses by priming CD4⁺ and CD8⁺ T cells.^{1,26,27} Upon recognition of PAMPs or DAMPs, DCs undergo a severe phenotypic and functional transformation, characterized by the secretion of a wide variety of inflammatory cytokines and the expression of maturation and activation markers such as MHC II and CD86. During this process, DCs become competent in cross-talking with lymphocytes, sustaining the expansion of specific effector cell subsets. As activation of the cGAS-STING pathway is well-known to promote DC activation, efforts have been made to understand the impact of STING activation in DCs within the tumor context. Accordingly, while it is widely accepted that DCs undergo a metabolic shift from oxidative phosphorylation to glycolysis upon activation, Hu et al recently found that increased ATP production via glycolysis facilitates STING-mediated antitumor functions in tumor-resident DCs.^{28,29} In an attempt to unravel the discrepancies observed with respect to the effects of STING activation in the tumor context in murine models and the human setting, Pang et al discovered that STING stimulation activated plasmacytoid (pDC) and conventional (cDC) DC subsets in both species, resulting in increased production of IFN I.³⁰ The authors also reported differential cell death responses upon STING activation, with rapid ablation occurring particularly for pDCs, highlighting the importance of understanding the impact of STING activation on different DC subsets.

cDCIs

cDC1s, identified by CD141⁺ in humans and CD103⁺ or CD8 α^+ markers in murine models, play a vital role in antitumor immunity by efficiently presenting tumor antigens on MHC I and thereby activating cytotoxic CD8⁺ T cells.^{1,31} Using

mice with deficiencies in different subsets of antigen-presenting cells, Wang et al demonstrated that specific STING licensing of cDC1s using cGAMP-loaded micelle nanoparticles (NPs) is essential for tumor rejection in mice. These NPs not only inhibited tumor growth in vivo but also induced a STING-activated cDC1 signature that correlated with higher survival rates among patients with non-small cell lung cancer (NSCLC).³² Another recent study found that the antitumor effects induced by cGAMP encapsulated in virus-like NPs also required STING activation in resident cDC1s (XCR1⁺ CD172⁻) but not in cDC2s. The choice of delivery vehicle, route of administration and STING agonist are all crucial factors, as the same study reported that administration of the synthetic CDN ADU-S100 induced cDC1 ablation.³³ In fact, ADU-S100 activates STING in cDC1s and not only fails to engage cDC2s but also induces upregulation of T-cell immunoglobulin and mucin-domain containing-3 (TIM-3), a traditional exhaustion marker.³⁴ Of note, another study reported that inhibiting TIM-3 also supports activation of the cGAS-STING pathway and increases the uptake of extracellular DNA in intratumoral cDC1s, emphasizing the role of TIM-3 as a regulator of DNA-sensing mechanisms.³⁵ Interestingly, cDC1 activation can also be prompted by IFN I secreted from STING-activated CD4⁺ T cells, thus establishing a mechanism that further supports antitumor immunity.³⁶

cDC2s

cDC2s, defined as CD1c⁺ and CD11b⁺ cells in humans and in murine animals, respectively, play a crucial role in halting cancer progression by directly promoting CD4⁺ T helper responses that support cytotoxic responses induced by CD8⁺ T cells.³⁷ Ulrich-Lewis et al convincingly demonstrated that upon treatment with a DNA vaccine, STING^{-/-} and cDC STING conditional knockout mice show lower Th1 responses compared to wild-type animals.³⁸ In line with previous findings, blocking of TIM-3 has been reported to enhance antitumor immunity mediated by STING agonist in 4T1 tumor models, by regulating cDC2s and improving the release of CD4⁺ T cells.³⁴

pDCs

Plasmacytoid DCs (pDCs) have a lower ability to act as antigen-presenting cells compared to cDCs. However, they are major producers of IFN and are key players in innate immunity, particularly in mounting responses against viral infections.^{39,40} In 2016, work by Bode et al demonstrated that human pDCs detect cytosolic DNA and CDNs via the cGAS-STING pathway, leading to robust activation and identifying pDCs as a potential target for new antitumor strategies.⁴¹ However, while stimulation of human pDCs with cGAMP or double-stranded immunostimulatory DNA (dsDNA) did induce nuclear translocation of IRF3, NF- κ B was not activated.⁴² Interestingly, STING signaling in pDCs has been shown to be susceptible to TME cues, as it is impaired in melanomas.⁴³ Furthermore, Brewitz et al reported that pDC-secreted IFN can support maturation and cross presentation of cDCs in close proximity, enhancing CD8⁺ T cell-mediated immunity, and thus pinpointing pDCs as potential targets of interest for novel cancer therapies.⁴⁴

Tumor-Associated DCs

In the context of cancer, all DC subsets may be present within the TME, albeit with different phenotypes and functions compared to DCs present in healthy tissues. Initially identified in NSCLC patients by Maier et al, tumor-associated DCs are characterized by high expression of activation markers such as MHC and CD80/86, as well as markers associated with immune tolerance such as indoleamine 2,3-dioxygenase (IDO) and programmed death-ligand 1 (PD-L1).^{27,45–47} Thus, tumor-associated DCs are often referred to as mature-regulatory DCs (mregDCs) and both type I and type II conventional DCs, cDC1s and cDC2s respectively, can acquire that phenotype. mregDCs are metabolically very active and play a critical role in modulating the TME by influencing the recruitment and function of other immune cells, exerting significant control over tumor progression.²⁷ By increasing the expression of regulatory genes, mregDCs can interfere with T-cell responses by favoring regulatory T-cell polarization and decreasing the number of antitumor effector T cells.^{48,49} Accordingly, while several studies describe tumor-suppressing effects of STING signaling, STING activation in the TME has also been reported to induce IDO expression in tumor-associated DCs in a murine model of lung carcinoma. Moreover, the authors showed that only in tumors with low antigenicity did DNA-sensing by STING promote a suppressive TME and thus tumor growth.⁴⁹ In contrast, activation of STING signaling in tumor-associated DCs in hepatocellular carcinoma models increased the secretion of CXCL9 and IL-12, leading to infiltration and activation of

natural killer (NK) cells.⁵⁰ The STING-mediated antitumor function of DCs can also be induced by oxidative stress in the TME. In fact, Hu and co-authors showed that oxidative stress is required for DC antitumor functions mediated by STING, as it promotes the accumulation of the redox sensor SUMO Specific Peptidase 3 (SENP3) which detects reactive oxygen species (ROS) and enables STING-mediated antitumor responses.⁵¹ Oxidative stress, mediated by ROS, is a feature of the TME that can activate immunosuppressive mechanisms and also induce DNA oxidation, which enhances immune recognition by DCs.^{33,34} Due to the rapid growth of cells during neoplasm formation, tumors are often deprived of oxygen, and the hypoxic TME can intensify the oxidative stress. This leads to a metabolic shift of tumor-resident DCs from oxidative phosphorylation to glycolysis, thus supporting downstream antitumor responses mediated by STING. As a result of STING activation induced by glycolysis, prompting DC activation.²⁹ Engagement of STING receptor is essential to start the activation and maturation of antigen-bearing tumor-resident DCs, supporting adaptive immunity by effector T cells (Figure 1).^{8,29,52}

STING Activation Strategies and Application in Cancer Therapies

Due to the strong immunostimulatory effect of STING signaling in immune oncology, much effort has focused on identifying the optimal agonist. Recent reports show that not only the induction of STING in immune cells can reduce TME-induced immunosuppression but also highlight that some agonists have the potential to enhance T cell infiltration at the tumor site, addressing a critical limitation of the success of adoptive cell therapy (ACT).^{53,54} Both Benoit-Lizon and



Figure I Illustration of the dual effects of STING activation in tumor-associated DCs. Tumor-associated DCs often display a mature-regulatory phenotype, characterized by the upregulation of both co-stimulatory as well as co-inhibitory molecules. In this context, it has been reported that the activation of STING in tumor-associated DCs may also play a dual role. On the one hand, engagement of the STING receptor can sustain antitumor effects by inducing HIF-1 α , which supports glycolysis in DCs, associated with an active and mature phenotype of these cells. This induces an IFN I-mediated response with secretion of pro-inflammatory cytokines that trigger effector T cell responses fighting the tumor (right panel). Conversely, activation of the STING pathway has also been reported to support pro-tumorigenesis, by inducing regulation of IDO expression and consequently enhancing regulatory T cell polarization (left panel). Created with BioRender.com.

Li have reported that STING agonists can directly activate T cells with no significant associated toxicity, improving effector responses mediated by CD8⁺ T cells, and Th1 and Th9, respectively.^{55,56} As STING induction may enhance antitumor immunity triggered by ICI and ACT, and as these therapies can be combined, it is possible that a suitable agonist may be the missing link for improved clinical success.^{56,57} Furthermore, Xu et al have shown that CAR T cells stimulated with STING agonists cGAMP or 5.6-dimethylxanthenone-4-acetic acid (DMXAA) improved the control of tumor development as it enhanced the permanence of CAR T cells at the tumor site. Of note, the authors also show that sustained anti-tumor responses required co-treatment with ICI or anti-GR-1 Ab, which deplete myeloid-derived suppressor cells.⁵⁴ The relevance of the cGAS-STING pathway as a druggable target in immune-oncology is largely acknowledged for its classical inflammatory response, but also for its ability to induce anti-cancer immunity via IFN I–independent mechanisms (described in¹⁰). Hines et al have nicely reviewed candidate agonist molecules currently being evaluated clinically.⁵⁸ In broad terms, STING-activating agents can be categorized into CDNs and non-CDNs of natural or of synthetic origin.

Natural Agonists in the Tumor Context

cGAMP is a CDN that is expressed in both physiological and pathological conditions.⁵⁹ It can activate STING directly in the cell of origin or it can act on surrounding cells.^{60,61} However, due to its hydrophilic nature, it cannot passively diffuse through the cell membrane, requiring protein transporters for intercellular access. Solute Carrier Family 19 Member 1 (SLC19A1) was the first reported cGAMP importer, but several others have been identified subsequently and were thoroughly reviewed by Blest et al.⁶⁰ Nevertheless, none of them is particularly efficient in mediating cGAMP internalization in primary cells, leading to very low accumulation of this agonist in DCs.^{60,62,63} Furthermore, cGAMP is rapidly destroyed by nucleases in the plasma, which limits its uptake by antigen-presenting cells under physiological conditions, thereby preventing constitutive STING activation and collateral chronic inflammation.^{64,65} In cancer, DCs are more exposed to STING agonists secreted by cancer cells in the TME because of chromosomal instability and DNA leakage. This is particularly notable in primary tumors, where cancer cell DNA-sensing mechanisms can autonomously trigger antitumor immunity. Consequently, byproducts of the cGAS-STING pathway such as cGAMP or dsDNA are fed into the TME through necrosis or cell leakage, boosting immunity via DC activation.^{7,66} In fact, recent studies have shown that the success rate of radiation as an anticancer therapy is dependent on STING signaling. Ionizing radiation may promote the activation of exosomes loaded with tumor-derived dsDNA that mediate the cargo transfer to DCs, initiating pro-inflammatory cascades.⁶⁷ This hypothesis is supported by another study where cancer cells were treated with another DNA damaging agent, a topoisomerase I inhibitor. This promoted the activation of DCs through delivery of tumor-associated DNA by exosomes.⁶⁸ Furthermore, ionizing radiation induces genotoxicity and immunogenic cell death not only in irradiated cells but also in neighboring cells, increasing the exposure of DCs to natural tumor-derived STING agonists.69-71

Synthetic Agonists

Synthetic agonists comprise CDNs and small molecules, often developed to improve the pharmacokinetics of STING agonists or to maximize STING signaling. The use of natural or synthetic CDNs as a standalone or combined cancer therapy is limited not only by their hydrophilicity, but also by their small size which causes rapid diffusion out of the tumors even when administered intratumorally.⁷² CDNs can be isolated from bacteria, such as cyclic di-GMP or cyclic di-AMP, or synthesized and modified to enhance their stability and immunogenicity for systemic administration.^{73,74} Despite achieving robust CD8⁺ T cell-mediated specific responses in preclinical models, prominent examples of synthetic CDNs like ADU-S100 did not perform well in clinical trials.^{75–78} Other CDNs have since been tested in clinical trials but all produced underwhelming results, as reviewed by others.^{79,80} To overcome the limitations of CDNs, other small molecules that sustain STING-mediated antitumor immune responses have been developed. For example, DMXAA was developed as an angiogenic disrupting agent in 2002, but it gained distinction for its potent immunomodulatory effects on the STING pathway, leading to unprecedented success in inducing tumor regression in pre-clinical models.^{81–83} However, DMXAA failed to replicate the results obtained in preclinical studies when used to treat NSCLC patients in combination with chemotherapy in Phase III trials. This disappointing result was traced to the drug's high specificity

exclusively for mouse STING.^{84–86} Several other non-CDN small molecules, most of which share structural similarities to DMXAA, are currently under development.^{80,87} For instance, dimeric amidobenzimidazole ligand (diABZI) is a synthetic non-CDN small molecule that demonstrated a 400-fold higher potency over cGAMP in activating peripheral blood mononuclear cells.⁶⁵ Nonetheless, small molecules lack strong cell selectivity across different subsets of immune cells, which leads to their uptake by lymphocytes and subsequent cell death. This highlights the need for drug delivery systems able to translate STING agonists into clinical applications.^{78,88–90}

STING Agonist Delivery Systems

Nano-based drug delivery systems offer promising advantages in modulating cargo pharmacokinetics and enabling targeted delivery, which can reduce drug-related side effects and enhance therapeutic effects.^{91,92} Liposomes were the first platforms investigated for encapsulating CDNs, due to their cationic lipid structure and aqueous core. In fact, encapsulation of cyclic di-GMP resulted in a STING-dependent enhanced antitumor response in metastatic melanoma mouse models.⁹³ Similarly, Doshi et al demonstrated that intravenous administration of ADU-S100 loaded into liposomes improved the therapeutic efficacy of that STING agonist and the responses to an immune checkpoint inhibitor (ICI) in both immunogenic and non-immunogenic in vivo tumor models.⁹⁴ CDNs were also successfully delivered using pegylated (PEG)-lipid discoid NPs, promoting efficient penetration into tumors after intravenous injection, and thus exposing most tumor cells to the loaded STING agonist in mice bearing colon adenocarcinomas.⁹⁵ The recently FDAapproved solid lipid nanoparticles (LNP) have also been exploited to this end, as they are known to successfully encapsulate nucleic acids and have intrinsic adjuvant properties. In line with this, Zhivaki et al have shown in vivo that LNPs loaded with mRNA encoding a self-DNA reactive variant of cGAMP had stronger immunostimulatory activity than other mRNA-encoded antigens also delivered by LNPs.⁹⁶ Mesoporous silica NPs also have been shown to be suitable for CDN delivery, achieving remarkable loading capacities and sustained release, resulting in a robust immune response in melanoma models.⁹⁷ Furthermore, several other polymeric-based NPs have successfully improved the delivery of CDNs, boosting the natural potency of cGAMP and inducing a "hot" immune TME in different tumor models.^{98,99} Accordingly, chitosan-based NPs, for instance, have been explored for their ability to induce cGAS-STING signaling, making them a promising adjuvant for use in novel formulations.^{100,101} Moreover, different ions, including Cu² ⁺, Zn²⁺, Mn²⁺, have been reported to support cGAS-STING signaling, but their exact mechanism has not yet been unveiled.^{102–105} In line with these findings, manganese has been deemed as essential not only in inducing cGAS-STING as a defense mechanism against dsDNA, but also in initiating innate immune responses to tumors by promoting maturation of DCs and macrophages and also supporting downstream effector T-cell responses.¹⁰⁶ The transporters for many of ions mentioned earlier are dysregulated in cancer (reviewed in¹⁰⁷), indicating that altered ion transport may be linked to cGAS-STING-mediated tumor immune evasion.

Nano-Based Platforms for Efficient Activation of STING in DCs

Several preclinical studies have explored different strategies to effectively deliver STING agonists to the TME and support STING-mediated tumor cell clearance. The interplay between cGAS-STING signaling and ions in the cancer context motivated the development of new strategies to tackle DCs with manganese nano-based systems.^{108–110} Using hybrid exosomes formulated with mannose ligands to target DCs, and protein nano-based structures without specific DC targeting, the effect of Mn²⁺ ions on STING activation of cDCs and bone marrow-derived DCs, respectively, was assessed in breast cancer tumor models. In both approaches, the authors included a DNA-damaging agent in their formulation that induces immunogenic cell death (ICD), thereby increasing the prevalence of dsDNA and conferring synergistic effects on STING engagement and further supporting antitumor responses mediated by the engagement of DCs with natural killer (NK) cells or with T-cells.^{111,112} Inducing tumor cell ICD to engage the cGAS-STING pathway has also been explored by Gu et al, in a study where colorectal cancer mouse models were co-treated with oxaliplatin-loaded liposomes and with ADU-S100.¹¹³ The authors observed that liposome-based immunotherapy not only induced cDC maturation, but also enhanced infiltration by CD4⁺ and CD8⁺ T cells. The effect of ADU-S100-loaded liposomes in DCs has also been assessed in colorectal and melanoma murine models. In this case, DCs were directly targeted by incorporating WH (a peptide with high affinity to Clec9a) into liposomes, and this resulted in enhanced potency of the

synthetic STING agonist.⁹⁴ The same DC-targeting strategy was used in another study, where poly(lactide-co-glycolide) (PLGA)-NPs encapsulating 2'3'-cGAMP were coated with another ligand of Clec9a, peptide CBP-12, and administered to animals with melanoma or breast cancer.¹¹⁴ The natural STING agonist 2'3'-cGAMP has also been encapsulated in virus-like particles and in liposomes and used to treat melanoma models. While there was no DC-targeting strategy for liposomes, in the virus-like particles a fusion glycoprotein was used for this purpose.^{33,115} To treat MC38-OVA tumorbearing mice, 2'3'-cGAMP was loaded into polymeric STING-activating NPs (PolySTING) and the antitumor effects observed were attributed specifically to cDC1 activation.³² A formulation with the same name, consisting of a prodrug platform that was used to deliver diABZI to DCs via mannose ligands, resulted in selective targeting of DCs in the TME and boosted not only cDC1/CD8⁺ T-cell crosstalk and lymphocyte infiltration, but also supported maturation of other DC subsets in tumor-draining lymph nodes.¹¹⁶ The recent observations made by Dosta et al, in which CDN-NP-mediated antitumor effects were assessed in melanoma models, further support the observations reported above. CDN-NPs behaved as a potent immune agonist, sustaining cDC activation either through direct uptake or through posterior transfer of NPs internalized by cancer cells.¹¹⁷ By exploiting natural intercellular mechanisms, Jang et al showed that DC targeting can be achieved to some extent by using CDN-loaded HEK293-derived extracellular vesicles (ExoSTING) and that this formulation can induce maturation of all DC subsets.¹¹⁸ To provide clarity on the efforts made to directly activate DCs in order to drive antitumor immunity, a summary of the effects of nano-based systems for activating the STING signaling pathway in DCs is provided in Table 1.

STING Controversy – the Tumor-Promoting Effect

Recent studies have highlighted the dual role of STING-induced responses in immune oncology. While activation of the STING pathway is crucial for natural antitumor immune responses in low immunogenic primary tumors, overactivation can lead to unproductive nonimmunogenic cell ablation, rendering patients more vulnerable to relapse or metastasis.^{33,78} In metastatic tumors with high chromosomal instability, targeting STING may further support tumor immune evasion.^{119,120} In fact, chronic activation of the cGAS-STING pathway, even at low levels, can sustain immunoregulatory gene expression such as IDO in DCs, thereby supporting tumor outgrowth and metastasis.^{49,121,122} Moreover, non-canonical activation of the STING pathway can also induce tumor-promoting responses, as such activation has been reported to support breast cancer cell survival in patient-derived xenograft models.¹²³ Persistent inflammation following STING engagement in phagocytic cells engulfing dead cells may be tumor-promoting, which is consistent with the theory that prolonged, unresolved inflammation contributes to oncogenesis. Thus, targeting the cGAS-STING pathway in DCs is particularly challenging, as DCs appear to waver between pro- and antitumor T-cell responses.

Concluding Remarks

The net effect of STING activation on cancer is heavily influenced by several factors, including the timing and magnitude of STING stimulation and the STING-responsive target cells. As this receptor is ubiquitously expressed in different cells and independent of pathological states, identification of an optimal STING agonist candidate has proven to be difficult, based on the results of clinical trials.^{80,90,124} Nevertheless, it is uniformly recognized that DCs play a crucial role in stimulating antitumor immune responses, with STING engagement being essential to prompt this process through IFN I secretion.¹⁰ However, because the development of DC-targeted vaccines for cancer therapy remains challenging, no standalone STING-based therapies have been successful to date.^{77,125} Instead, current approaches combine the administration of STING agonists with ICIs such as PD-L1 or CTLA-4, because at optimal doses, STING induction prompts antitumor immunity mediated by IFN I signaling and it facilitates tumor infiltration by T cells, thereby converting "cold" tumors into "hot" tumors that are more prone to respond to ICI therapies.^{8,9,126–129} However, the underwhelming results of this approach underline the challenge of applying STING agonists in the clinic. The major hurdles hindering the clinical success of STING engagement in cancer immunotherapy are not only the poor pharmacokinetic and physico-chemical properties of CDNs that limit their systemic administration, but also the frequent impairment of STING signaling in multiple cancers, although its restoration may be achieved by epigenetic reprogramming.^{80,130–132} To overcome these difficulties, small molecule STING agonists have been tested in combination with immune checkpoint

Table I Strategies for Activation of the STING Pathway in DCs

Delivery System	Cargo	Size (nm)	DC-targeting Approach	DC Subset Analyzed	Model	Route of Administration	Ref
Hybrid exosomes	SN38 and MnO2	149	Unspecific (ICD of cancer cells and stimulation of STING)	cDC (CDHc⁺)	4TI tumor model	i.v.	[112]
Protein-based nanostructure	Mn2+ and β -lapachone	100	Mannose ligands	BMDC	4TI tumor model	i.v.	[11]
(Man@Mn2+-Ft@Lap) Liposomes	Oxaliplatin	122	Unspecific (ICD of cancer cells and stimulation of STING through co- administration of ADU-S100)	cDCI and cDC2 (CDIIc⁺)	CT26 murine model	Liposomes: i.v. ADU-S100: i.t.	[113]
Liposomes	ADU-SI00	121.3	WH peptide (Clec9a targeting)	cDCI (CDI03⁺)	MC38 and B16F10 murine	i.v.	[94]
PLGA NPs	2'3'-cGAMP	157.5	CBP-12 EPBM NP coating (Clec9a targeting)	cDCI (CD8α ⁺)	tumor BI6-OVA melanoma and 4TI breast cancer	s.c.	[114]
Liposomes	2'3'-cGAMP	160	Unspecific	BMDC	BI6-FI0 melanoma model	in vitro stimulation	[115]
Virus-like particles	2'3'-cGAMP	158	Viral fusion glycoprotein	cDCI and cDC2 (CDIIc ⁺)	B16-F10 melanoma model	i.t.	[33]
PolySTING	2'3'-cGAMP	25	Not specified	cDCI (CD8 α^+ and CD103 ⁺)	MC38-OVA tumor- bearing	i.t.	[32]
PolySTING	diABZI	-	Mannose ligands	cDCI (CD8 α^+ and CD103 ⁺)	BI6FI0 and 4TI tumor models	i.v.	[116]
CDN-NPs	Synthetic CDN	41	Unspecific	cDC (CDIIc⁺)	B16-F10 melanoma model	i.v.	[117]
ExoSTING	MR SS-2 CDA or cAIM(PS)2 Difluor (Rp/Sp)	50–200	EV import	pDC (CD123 ⁺), cDC (CD11c ⁺)	B16F10, EG7.OVA and CT26 models	i.t.	[118]

Abbreviations: BMDC, bone marrow-derived dendritic cell; i.v., intravenously; i.t., intratumorally; s.c., subcutaneous.

blockade, but these attempts were reported to induce toxicity in lymphocytes, highlighting that antitumor immunity requires STING licensing specifically in myeloid cells.^{88,133}

In summary, the cGAS-STING pathway is an undisputed target for novel developments in cancer immunotherapy, mediating the orchestration of critical antitumor responses by tumor-resident DCs. However, although STING signaling is essential to combat cancer progression, recent evidence has revealed its dual role, where chronic activation can paradoxically support tumor growth. Thus, the future of immuno-oncology relies on a deeper understanding of the effects of STING activation on different DC subsets in the tumor context, as well as on the development of efficient, tailored STING agonist delivery systems that will allow for optimal antitumor immune responses while mitigating potential side effects.

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Disclosure

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