IDO in the Tumor Microenvironment: Inflammation, Counter-regulation and Tolerance

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Abstract

Indoleamine 2,3-dioxygenase (IDO) has immunoregulatory roles associated tryptophan metabolism. These include counter-regulation (controlling inflammation) and acquired tolerance in T cells. Recent findings reveal that IDO can be triggered by innate responses during tumorigenesis, and also by attempted T cell activation, either spontaneous or due to immunotherapy. Here we review the current understanding of mechanisms by which IDO participates in the control of inflammation and in peripheral tolerance. Focusing on the tumor microenvironment, we examine the role of IDO in response to apoptotic cells and the impact of IDO on Treg cell function. We discuss how the counter-regulatory and tolerogenic functions of IDO can be targeted for cancer immunotherapy and present an overview of the current clinical progress in this area.

Introduction

The biologic function of the IDO pathway is both counter-regulatory (controlling inflammation) and tolerogenic (creating acquired antigen-specific tolerance in T cells) [1]. Thus, when IDO is genetically deleted or pharmacologically inhibited, mice experience defects in creating tolerance to the fetus during allogeneic pregnancy [1-3], mucosal tolerance [4,5], tolerance to apoptotic cells [6,7] and other forms of acquired peripheral tolerance [8-13]. Conversely, gene-transfer or pharmacologic induction of IDO can create de novo systemic tolerance [6,13-15]. IDO also enhances the tissue-reparative effects of inflammation while limiting the tissue-destructive effects. Thus, blocking or ablating IDO makes inflammation markedly worse in models of graft-versus-host disease [16-17], autoimmunity [18-21] or chronic infection [22-23]. In all of these models, however, the role of IDO is focused and selective, and IDO-deficient mice do not have the broad, spontaneous defects in self-tolerance that are seen with mice lacking CTLA-4 or Treg cells. But in...
settings where IDO is crucial, it can be an important and non-redundant mechanism of immune regulation.

IDO creates its biologic effects by metabolizing the essential amino acid tryptophan into kynurenine. The IDO family of genes includes IDO1 (the main subject of this review) and IDO2 [24–26]. IDO2 is not redundant with IDO1 but it is much less well studied. In this review, the term “IDO” will refer to IDO1, or to collective functional IDO enzyme activity, unless otherwise specified. Depletion of local tryptophan by IDO can activate the stress-response kinase GCN2 [27–28], which senses amino acid withdrawal. GCN2 activation in T cells can inhibit their proliferation, and can bias naive CD4+ T cells toward differentiation into Treg cells [27–29]. In addition, IDO produces soluble factors (kynurenine and downstream metabolites) that bind and activate the aryl hydrocarbon receptor (AhR) [30]. The AhR can promote Treg cell differentiation [29–30], and can also bias dendritic cells (DCs) and macrophages toward an immunosuppressive phenotype [6·31–33]. GCN2 can also directly affect the phenotype of dendritic cells and macrophages [6·33–34]. Together, these pathways can have a profound effect on the APC and the antigen-presenting milieu. Thus, when IDO is active, APCs otherwise capable of producing inflammatory cytokines such as IL-12 instead produce inhibitory cytokines such as IL-10 and TGFβ [6·7·34]. This is a key concept, because it means that IDO up-regulation can alter the nature of the APC itself, and change the whole local milieu from immunogenic to tolerogenic [6·35].

In 2013 in this journal, we reviewed the role of IDO in creating metabolic signals that regulate immune responses [36]. We discussed the ability of IDO to create local suppression of effector T cells by metabolic depletion of tryptophan and production of tryptophan metabolites; and also to create more widespread systemic tolerance by activating circulating Tregs. We also noted the counter-regulatory role of IDO, which is actively induced by many of the same inflammatory signals that it acts to control. Since that time, IDO has entered multiple Phase II clinical trials in oncology, and has become an increasingly attractive target for cancer immunotherapy. IDO has been shown to play an important role in inducing tolerance to apoptotic cells, which is relevant both for self-tolerance and for the immune response to dying tumor cells. In the current review, we examine how counter-regulatory IDO can be triggered in the tumor microenvironment by innate responses during tumorigenesis, and in established tumors by attempted T cell activation (either spontaneous or due to immunotherapy). We will describe how IDO may affect signaling pathways in Tregs via PTEN phosphatase, mTOR and PI(3)K. Overall, IDO possesses somewhat different properties from other immune checkpoints in the tumor microenvironment, because it can act upstream to affect the basic choice between tolerogenic or immunogenic cross-presentation of tumor antigens.

**IDO and counter-regulation**

The tumor microenvironment is an abnormal milieu, but tumors often recruit and exaggerate the normal regulatory mechanisms that maintain homeostasis and tolerance in the immune system. As we will discuss, many tumors over-express IDO, either in the tumor cells themselves, or in tumor-associated cells such as dendritic cells (DCs), macrophages or endothelial cells. Further complicating this biology, IDO is inducible, and may not be
expressed until some degree of inflammation occurs in the tumor to drive it [37]. This is a key point, because IDO by its nature tends to be counter-regulatory: i.e., its expression is actively induced by signals from inflammation or T cell activation, which it then acts to suppress. This inducible counter-regulation is beneficial when IDO is controlling harmful inflammation or creating tolerance to apoptotic cells, but is highly undesirable when it is suppressing the immune system’s attempted response against the tumor.

**Physiologic IDO induction at sites of inflammation**

Interferon types I (IFNαβ) and II (IFNγ) are classic hallmarks of inflammation that stimulate immunity. IFNs activate Janus kinase/signal transducer and activator of transcription (JAK/STAT) complexes to induce transcription of many IFN-stimulated genes (ISGs). Many ISGs promote immunity, but some ISGs regulate immunity and promote tolerance. IFNs are potent inducers of immune regulatory responses mediated by IDO. Mammalian IDO1 gene promoters possess IFN-stimulated response elements (ISRE) and IFN-activated sites (GAS). In contrast, IDO2 genes are weakly responsive to IFNs. Other factors that induce IDO in some cells include aryl hydrocarbon receptor (AhR) ligands and regulatory cytokines, including IL-10 and TGFβ. Select cell types express IDO at sites of inflammation, even though most cells express IFN type I (IFNAR) and type II (IFNγR) receptors. Infections, vaccines and tumorigenesis induce IFNs that may co-induce IDO, though IDO enzyme activity may not manifest in these settings due to transcriptional, translational and post-translational controls. Some epithelial cells, dendritic cells (DC) and macrophages can express IDO, while lymphoid cells rarely express IDO. The transcriptional factor DAP12 regulates IFN-induced IDO transcription while SOCS3 targets IDO protein for proteosomal degradation in some cells [38-39]. Metabolic factors that regulate IDO enzyme activity include heme cofactor and substrate supply and redox potential. Cells expressing IFNγ-inducible nitric oxide synthase (iNOS) produce nitric oxide (NO) that blocks IDO activity. Hence IFNγ co-induces iNOS and IDO but metabolic cross-regulation may override IDO-mediated. In summary, many inflammatory insults induce IDO but multiple factors restrict IDO expression and regulate IDO activity in physiologic settings. IDO expression in tumors

In some tumors, IDO is constitutively expressed by the tumor cells themselves. This may serve as an immune-escape mechanism (which is the focus of this review), or in some cases may confer some non-immune survival advantage on the tumors [40-41]. IDO can also be expressed in tumor-associated cells such as DCs or endothelial cells. In most studies, high expression of IDO in the tumor or draining LNs has been an adverse prognostic factor. Tumors in this category include melanoma [42-44], colon cancer [45-46], brain tumors [47], ovarian cancer [48], acute myelogenous leukemia [49-50], and a number of others [51-53]. In a smaller number of tumor types, IDO expression appears to be induced or “reactive” – i.e., associated with increased T cell infiltration and inflammation [37-54-55]. In this situation, up-regulation of IDO may be a proxy for a stronger spontaneous anti-tumor immune response, and thus associated with more favorable prognosis [56-57]. However, even in these immune-responsive patients, the IDO itself is not beneficial, and the patient might do even better if IDO were blocked (this will need to be evaluated empirically in the course of clinical trials). The signals that upregulate reactive IDO in these tumors are not
known, but IFNγ would be a logical candidate, and there is experimental support for this hypothesis [37].

Growing awareness that beneficial anti-tumor inflammation can induce detrimental counter-regulatory IDO has implications for clinical immunotherapy. The goal of most immunotherapy is to activate T cells within the tumor. If these T cells themselves induce IDO, this might reduce or inhibit the desired anti-tumor effect. While this concern has not yet been tested in clinical trials, preclinical examples have been demonstrated with IL-12 therapy [58] and adoptive cell therapy using CAR-T cells [59]. Thus, immunotherapy regimens that induce extensive inflammation at the site of the tumor might benefit from combination with an IDO-inhibitor drug.

**Immune responses to DNA in the tumor microenvironment**

DNA is sensed to activate immune cells. This response is critical for host control of many infectious pathogens, including bacteria ingested by phagocytes and DNA viruses that replicate in cells. Host survival may depend on rapid induction of innate immunity to limit infection spread and incite adaptive immunity to eliminate infections. Toll-Like Receptor-9 (TLR9) is the archetypal DNA sensor that recognizes un-methylated CpG motifs in prokaryotic (bacterial) DNA and mitochondrial DNA. TLR9 expression is restricted to specialized immune cells, including some DCs, macrophages and B cells. These cell types function as sentinels and antigen presenting cells (APCs) that respond rapidly to infections and provoke host immunity. TLR9 is located in endosomes, where it senses ingested pathogen DNA to induce IFNαβ expression via MyD88 signaling, a classic hallmark of host innate immunity to microbial infection.

Eukaryotic cells also express the signaling adaptor STimulator of INterferon Genes (STING, also termed MPYS, Tmem173). STING is downstream of several cytosolic DNA sensors [60], including cyclic GMP-AMP synthase (cGAS). STING interacts with TBK1 and IRF3 signaling proteins and IFNβ is a prominent target of STING signaling. The mechanism by which DNA sensing activates STING is known for cGAS, which possesses ‘zinc thumb’ structures that bind to double strand DNA (dsDNA). Once bound to DNA, cGAS undergoes conformational changes that catalyze synthesis of the cyclic dinucleotide (CDN) cGAMP [61]. Cytosolic DNA sensing by cGAS to activate STING/IFNβ signaling are key steps in rapid responses to infection by many professional APCs and non-APCs.

**Cytosolic DNA sensing can incite tumor immunity**

STING/IFNαβ signaling also incites tumor immunity in mouse models of spontaneous or therapy-induced tumor regression [62–63]. Thus B16 melanomas expressing a neo-antigen (SIY) grew faster in STING-deficient mice than STING-sufficient (B6) mice. DNA was sensed via cGAS to activate STING/IFNαβ signaling and stimulate tumor antigen cross-presentation by CD8α+ DCs that primed T cells. Hence in this immunogenic B16-SIY tumor model, STING/IFNαβ signaling led to dominant pro-inflammatory responses that impeded tumorigenesis (Figure 1A). These observations identified DNA as a source of pro-inflammatory signals that incite tumor immunity in the context of immunogenic tumor
growth. Malignant cells or tumor-associated cells that die are potential sources of DNA that trigger STING/IFNαβ signaling in the tumor microenvironment. Phagocytes in tumor lesions may ingest dying cells and process DNA and tumor antigens to prime T cells. Strong correlations between IFNαβ transcriptional signatures and T cell markers in human metastatic melanomas provide clinical significance for the paradigm that DNA sensing in the tumor microenvironment incites T cell responses [64]. This paradigm also suggests that STING agonists may be exploited to enhance tumor immunity. Indeed, the tumor-disrupting agent DMXAA (Vadimezan) and modified CDNs that activate STING were potent adjuvants for tumor immunotherapy in mice [65]. However, therapeutic responses to Vadimezan were not IFNαβ-dependent, hinting that other pathways may incite tumor immunity downstream of STING. Clinical trials revealed no therapeutic benefit of Vadimezan, possibly because human STING is not responsive to the drug due to an amino acid polymorphism [66]. This point notwithstanding, modified CDNs that activate human STING may be effective adjuvants in some cancer patients based on promising therapeutic responses observed in mouse immunogenic tumor models [65].

DNA and STING agonists can promote tolerogenic responses by inducing IDO

DNA nanoparticles (DNPs) containing cargo DNA and cationic polymers such as polyethylenimine (PEI) are used to transfect genes into cells and live organisms. Unexpectedly, systemic DNP treatments stimulated IDO activity in many mouse tissues due to cargo DNA sensing by innate immune cells [67]. Cargo DNA lacking TLR9 ligands was sensed to induce IDO via a STING/IFNαβ dependent pathway, though prokaryotic DNA was also sensed via TLR9 to activate NK cells [35]. Cargo DNA was ingested and sensed via STING to stimulate selective IFNβ expression by myeloid (CD11b) DCs and induce selective IDO up-regulation in DCs expressing the B cell marker CD19. Splenic DCs from DNP-treated mice suppressed T cell responses, activated Tregs and DC ablation abolished IDO-mediated suppression following DNP treatment. Thus discrete DC subsets in lymphoid tissues sense cytosolic DNA to activate STING and mediate IDO-dependent tolerogenic responses.

Consistent with the paradigm that STING can regulate immunity, STING agonists alleviated autoimmune disease syndromes in mice. Systemic DNP treatments alleviated limb joint damage in an antigen-induced arthritis model [67], while DNPs and synthetic CDNs slowed experimental autoimmune encephalitis (EAE), a model of multiple sclerosis, and also alleviated established EAE [68]. In both models, tolerogenic and therapeutic responses to STING agonists were abolished in mice lacking STING, IFNαβ receptor or IDO1 genes, and in mice given IDO inhibitor. Thus when administered systemically STING agonists are tolerogens due to their ability to induce IDO.
Cellular DNA in aseptic tissues is sensed to incite or inhibit autoimmunity via STING

Diametric immune responses to cytosolic DNA during immunogenic tumor growth and after DNP treatment highlight the critical need to understand how cellular DNA is processed and sensed to activate STING in local settings of inflammation. Release of DNA and intracellular autoantigens from dying cells may compromise tolerogenic processes that prevent autoimmunity. Moreover, inflammation lowers tolerance thresholds and increases cell death to enhance the risk of inciting autoimmunity. Unlike live cells that incite immunity, apoptotic cells promote tolerance to alloantigens they express. Systemic administration of apoptotic thymocytes incited robust tolerogenic responses in mice via an IDO-dependent pathway [7]. Thus pre-treating female B6 mice with male apoptotic cells protected male skin allografts from rejection, while male allografts were rejected by females given IDO inhibitors during pre-treatment [6]. Moreover, STING ablation abolished IDO and regulatory cytokine (IL-10, TGFβ) induction after apoptotic cell treatment, which instead stimulated pro-inflammatory IL-6 expression [35]. Thus apoptotic cell DNA was sensed via cytosolic DNA sensors to promote tolerance via IDO. Like DNPs that resemble chromatin, DNA complexes from dying cells may be ingested by phagocytes that process DNA to activate STING/IFNαβ signaling and promote tolerogenic responses via IDO. Marginal zone (MZ) macrophages expressing CD169 mediated tolerogenic responses to apoptotic cells, though other MZ cell subsets may collaborate with CD169+ MZ to coordinate processing of apoptotic cell antigens and DNA to elaborate tolerogenic responses. By analogy, immune cells that infiltrate developing malignancies or reside in draining lymph nodes may be specialized to promote tolerance via IDO in response to cytosolic DNA sensing. Some potential factors likely to influence immune responses to DNA in the tumor microenvironment include: (1) cell death rates and pathways of cell death; (2) physiologic context that may promote distinct inflammatory responses during tumorigenesis and; (3) genetic, environmental and immunologic factors that impact the local balance between tolerance and immunity.

Striking illustrations of the potential for tissue DNA to break tolerance emerged from studies on mice with defects in DNA catabolism. Mice lacking the DNAses Trex-1 or DNAse-II succumbed to spontaneous hyper-inflammatory and autoimmune syndromes, leading to death several months after birth or embryonic death, respectively [69, 70]. In these models, spontaneous autoimmunity was caused by excessive tissue DNA sensing to activate STING and incite sustained IFNβ production. Thus in aseptic tissues DNA incited autoimmunity unless DNAse activity degraded excess DNA, suggesting that cell death during tissue homeostasis, growth or re-modeling has the potential to incite STING/IFNαβ signaling that can overcome local tolerance.

Conversely in some physiologic settings, active tolerogenic responses to tissue DNA may maintain peripheral tolerance when cells die, even when DNAse activity is not compromised. Support for this paradigm emerged from studies on MRL<sup>lpr</sup> mice prone to systemic lupus-like autoimmune syndromes since STING-deficient MRL<sup>lpr</sup> mice were more susceptible to lupus than their STING-sufficient counterparts [71]. Moreover, macrophages...
were hyper-responsive to TLR ligands, while IDO expression and Treg numbers were reduced substantially in STING-deficient MRL<sup>lpr</sup> mice. These observations are consistent with findings that IDO inhibitors accelerated lupus progression in MRL<sup>lpr</sup> mice and that IDO1 gene ablation conferred lupus susceptibility following chronic exposure to apoptotic cells [7]. Therapeutic responses to STING agonists that alleviated autoimmune diseases also support the paradigm that DNA sensing to activate STING can reinforce tolerogenic processes [67–68]. Collectively, studies on mice with defective DNase function and MRL<sup>lpr</sup> mice reveal opposing immunologic responses to cytosolic DNA that incite or suppress autoimmunity, respectively. Thus cytosolic DNA sensing to stimulate STING/IFN<sub>αβ</sub> signaling is a pivotal pathway able to drive dominant tolerogenic or immunogenic responses. Multiple factors are likely to impact the direction of immune responses to DNA, STING agonists and dying cells. For example, systemic administration of DNPs, STING agonists and apoptotic cells is essential for dominant tolerogenic responses to manifest in mice.

**STING and IDO in the tumor microenvironment**

The paradigm that DNA can drive opposing immunologic outcomes is pertinent to understanding how DNA influences immune responses in the tumor microenvironment. As discussed above, DNA sensing to activate STING/IFN<sub>αβ</sub> signaling slowed growth of immunogenic tumors and potentiated tumor regression after therapy [63–72,74]. Moreover, IFN<sub>αβ</sub> signaling impeded growth of carcinogen-induced and transplantable tumors [75]. It is presently unclear how immunogenic responses driven by IFN<sub>αβ</sub> can be reconciled with the paradigm that IDO, which is up-regulated by IFN<sub>αβ</sub>, is often up-regulated during tumorigenesis in mice and cancer patients. Factors that induce and sustain IDO prior to overt tumor formation have not been defined, but may include DNA (by analogy to physiologic IDO induction following DNP or apoptotic cell treatments). Elevated IDO may be an early ‘innate’ response to local inflammation associated with pre-malignant lesions. IDO may also be induced at later stages of tumorigenesis, when activated myeloid (DCs, Mϕs, MDSCs,) or lymphoid (T, NK) cells infiltrate tumor lesions. IFNs are likely inducers of IDO, though IFN<sub>αβ</sub> may be more likely to induce IDO during the early stages of tumorigenesis when cells making IFN<sub>γ</sub> are absent. IDO1-deficient mice were resistant to skin papilloma formation in the DMBA/TPA carcinogenesis model [76] suggesting that IFNs may induce IDO to promote immune escape in this model. Moreover, mice lacking STING genes in bone marrow derived cells were also resistant to papilloma formation after chronic DMBA exposure [77], revealing a critical role for cytosolic DNA sensing by hematopoietic cells in carcinogenesis. These observations suggest that DNA enters cytosolic compartments of hematopoietic cells to promote carcinogenesis, possibly due to STING/IFN<sub>αβ</sub> signaling that induces IDO. Infiltrating effector CD8 T cells were also identified as sources of IFN<sub>γ</sub> that induced IDO and create immune checkpoints in the tumor microenvironment of malignant melanomas [37].

Another factor that may influence immune responses to DNA in the developing tumor microenvironment is tumor antigenicity, which is gaining recognition as a key factor determining responsiveness to immunotherapy [78]. As noted above, immunogenic B16-SIY melanomas grew faster in STING-deficient mice. In the same study, parental B16 melanomas grew at comparable rates in STING-deficient and wild-type mice not given
therapy [65]. Thus DNA sensing to activate STING had no significant impact of
tumorigenesis in the absence of tumor neo-antigens, indicating that enhanced tumor antigen
recognition was essential for DNA to promote tumor immunity, lower tolerance thresholds
and impede tumorigenesis (Fig. 1A). Tumors with relatively low antigenicity may be less
likely to overcome innate tolerizing responses to DNA that suppress tumor immunity and
promote tumorigenesis (Fig. 1B). Similar questions arise regarding immune responses to
DNA following therapy because DNA sensing may help restore or reinforce tumor tolerance
by stimulating IDO. Reports that STING impeded growth of immunogenic tumors and
mediated therapeutic effects suggest that immunogenic responses to DNA overcame local
tolerogenic processes in these models [62-63]. However this paradigm may not apply to all
tumor models, especially tumors with low antigenicity. In such settings tolerance may be
more robust, making it more difficult to overcome counter-regulatory responses established
during tumorigenesis, including DNA driven IDO up-regulation (Fig. 1B). Similar
considerations may also help explain why therapeutic responses to STING agonists (CDNs)
were effective in some but not all tumor models [63-65]. Thus, DNA sensed in the tumor
microenvironment creates an innate immune adjuvant effect that can potentiate efforts to
stimulate tumor immunity. However, immune responses to DNA during tumorigenesis and
in response to immunotherapy can vary – boosting immune recognition and effector
responses, or inducing tolerogenic IDO, as it does for DNP cargo DNA and DNA released
by apoptotic cells. Thus, it may be necessary to determine empirically whether DNA sensing
to activate STING promotes immunity or tolerance in a particular tumor setting. This point
notwithstanding, it seems likely from recent developments that STING agonists may be of
most benefit in tumors that exhibit relatively high immunogenicity.

**IDO and Treg cells**

We now turn from innate inflammation to regulation of adaptive immune responses. One of
the major pathways by which IDO can affect T cell immune responses is via activation of
suppressive Treg cells.

**Activation of Treg cells by IDO**

Regulatory T cells are a major suppressive mechanism in tumors [79]. IDO controls one
important activation pathway for Tregs in the tumor microenvironment. Resting Tregs are
not suppressive [80] and must undergo some form of activation in order to become
functional. This activation step requires TCR engagement [81], but it is also highly sensitive
to modulating signals from the local milieu. These environmental signals can dictate the
functional properties of the activated Tregs [82]. When Tregs are activated by an IDO-
expressing APC, they acquire properties that are very different from the conventional in
vitro systems using anti-CD3 mitogen [83]. One difference is that IDO-induced Treg
activation is triggered by physiologic signals from activating effector T cells; this has
recently been shown to be a key stimulus for Treg activation in vivo [84]. A second
important difference is that the presence of IDO markedly affects the outcome of Treg
activation by regulating mTORC2 and Akt signaling in the Treg, as summarized in Figure 2.
One of the effects of IDO is to reduce the local tryptophan concentration in the vicinity of the responding Treg. This low tryptophan activates a stress-response pathway in the Tregs mediated by the kinase GCN2 [27–85]. Then, via a process that is not yet well understood, GCN2 inhibits the activity of the mTORC2 complex, and prevents it from phosphorylating Akt on its activating Ser473 site [86]. Akt is emerging as an important control-point for Treg activation in tumors [86–87]. Unlike effector T cells, which need Akt signaling for normal activation, Akt signaling in Tregs inhibits their function [88]. More precisely, Akt destabilizes the Tregs and causes them to lose their suppressive activity [86–89]. These destabilized Tregs may then reprogram into pro-inflammatory effector cells [90–92], which have been termed “ex-Tregs” [89–93–94]. Thus, one of the important functions of IDO during Treg activation is to inhibit Akt and thus prevent destabilization and maintain the suppressive phenotype.

Akt triggers inactivation and degradation of the FOXO transcription factors FoxO1 and FoxO3, which are important for Treg suppressor function [95]. Thus, by inhibiting the mTOR/Akt axis, IDO allows Tregs to up-regulate FoxO3a; and this in turn allows them to up-regulate PD-1 expression [86]. When this PD-1 is engaged by its ligands PD-L1 or PD-L2, it activates the PTEN lipid phosphatase, which then inhibits PI3K activity and blocks phosphorylation of Akt on its other activating site, Thr308 (Figure 2). Thus, IDO-activated Tregs establish a PD-1→PTEN signaling loop that stably maintains inhibition of Akt, as long as PD-1 is engaged by its ligands. (And, since IDO-activated Tregs are potent inducers of PD-L1 and PD-L2 expression on DCs [83], it is likely that PD-1 will remain engaged.) This PD1→PTEN feedback loop thus provides a molecular explanation for the earlier observation, seen in multiple models, that IDO-induced Treg activity is strictly dependent on the PD-1/PD-ligand pathway in order to create suppression [67–83–96–97].

Taken together, these data suggest that when Tregs are activated in the presence of IDO, this initiates a stable self-perpetuating suppressive state in the Tregs that becomes independent of IDO, and instead is maintained long-term by the loop between PD-1 signaling and PTEN. This has important implications, because IDO expression is localized and relatively restricted, whereas PD-ligands are widely inducible on antigen-presenting cells, tumor cells and other tissues. Thus, IDO-activated Tregs could potentially create sustained suppression far from the original source of IDO. More speculatively, it is possible that other metabolic stresses that could block the mTOR/Akt pathway during Treg activation (e.g., arginine deprivation due to arginase I expression) might also initiate the same stable, PTEN-driven activation loop. Conversely, activation signals might target PTEN directly. Neuropilin-1 (Nrp1) on Tregs is an important pathway that contributes to immune suppression within the tumor microenvironment [98]. Nrp1 activates PTEN, inhibits Akt, preserves FoxO3a and stabilizes Treg suppressor activity [98]. Thus, Nrp1 might establish the same stable, self-sustaining, PTEN-driven Treg phenotype via direct action on PTEN, and IDO may be just one of several upstream pathways that converge on PTEN during Treg activation.
IDO and tolerance to apoptotic cells

Physiologic tolerance to apoptotic self cells

One important physiologic pathway that may depend strongly on the link between IDO and PTEN is the tolerogenic response to apoptotic cells. IDO appears to play a significant role in maintaining tolerance to apoptotic cells. When normal mice are challenged with apoptotic thymocytes, they maintain tolerance to self antigens [7], and will create de novo tolerance to new antigens introduced on the apoptotic cells [6]. However, if IDO1 is genetically ablated or pharmacologically inhibited, then mice can no longer maintain tolerance when challenged with apoptotic cells, and rapidly develop lupus-like autoimmunity [6, 7]. Apoptotic cells injected intravenously were taken up by macrophages in the spleen, which triggered up-regulation of IDO [7]. IDO then altered the phenotype of both the macrophages and of neighboring cells: promoting expression of tolerogenic IL-10 and TGFβ, inhibiting production of immunogenic IL-12, altering the phenotype of local cross-presenting DCs, and recruiting suppressive Tregs [6, 99]. Thus, IDO affected more than just T cell activation: it modified the properties of the antigen-presenting cells themselves, and altered the whole antigen-presenting milieu. Relevant to tumor immunology, the key point from these studies is that apoptotic cells were not inherently “invisible” or tolerogenic. If they were prevented from inducing IDO, then even self cells became inflammatory and immunogenic.

In the case of apoptotic tumor cells, a similar tolerogenic role for IDO was observed. Subcutaneous injection of apoptotic tumor cells caused activation of IDO, with IDO-dependent induction of suppressive PTEN-expressing Tregs [86]. Mice with a genetic deletion of PTEN in Tregs could not create IDO-induced tolerance to the apoptotic tumor cells [86]. (Of note, these PTEN-deficient mice also show the same lupus-prone phenotype as IDO1-KO mice when challenged with apoptotic self cells [86].) Generalization of these findings to the tumor microenvironment in vivo is still speculative, but tumors are constantly faced with the need to inhibit the immune response to dying tumor cells. This becomes particularly acute during the wave of tumor cell death after chemotherapy or radiation. Thus, given the natural physiologic role of IDO in eliciting tolerance to apoptotic cells, it may be that tumors “hijack” this pathway to suppress immune response against dying tumor cells.

IDO and immunogenic tumor cell death

In principle, tumor cells should be much more immunogenic than normal cells. Tumors have many mutational neoantigens, and even authentic self antigens may be aberrantly expressed in such a way that they become immunogenic [100]. Consistent with this, certain forms of chemotherapy have been shown to elicit a form of immunogenic tumor cell death that triggers inflammation and T cell responses (reviewed in ref. [101]). Given the widespread use of chemotherapy in the clinic, this is potentially an extremely valuable pathway. However, in practice this phenomenon has been restricted primarily to anthracycline chemotherapy, and to certain transplantable tumor models. But if it could be made more widely applicable, immunogenic cell death could be a powerful tool for therapy. Hence, there is considerable interest in enhancing the immunogenic effects of chemotherapy by combination with immunomodulatory agents [102-103].
The role of IDO and IDO-activated PTEN-Tregs in suppressing the response to apoptotic cells raises the possibility that immunogenic cell death may actually be more widespread – and occur in more settings – than previously recognized. Figure 3 summarizes the possibility that, during tumor cell death, two sets of opposing signals may be created simultaneously. When tumor cells die – whether by chemotherapy, radiation or immunotherapy – some fraction will presumably die by traditional apoptosis. These apoptotic tumor cells will deliver the same tolerogenic signals created by any apoptotic cells (TGFβ, induction of IDO via STING and other pathways, and IDO-induced activation of PTEN-Tregs), as described in refs. [7] and [86]. Simultaneously, however, any tumor cells that die via dysregulated forms of cell death (immunogenic cell death, necrosis or necroptosis) will deliver inflammatory signals. Since tumor cells are so aberrant and disorganized, it seems likely that there may always some component of immunogenic (inflammatory) death combined with the apoptotic cell death. The question then becomes which set of signals – tolerogenic or immunogenic – is dominant. Unless the signal from immunogenic cell death is very strong, IDO and Tregs may predominate – thus masking the fact that many tumor cells are potentially immunogenic. This hypothesis is at present still speculative. If correct, however, it would mean that immunogenic tumor cell death may already be widely available in many contexts (e.g., with standard treatments in the clinic), if the dominant effects of IDO and the downstream PTEN-Treg pathway can be removed.

Interaction between IDO and other pathways in the tumor microenvironment

IDO and other checkpoints

Currently, inhibitors of the CTLA-4 and PD-1 pathways are rapidly advancing in the clinic [104]. Results are exciting for certain types of tumors, especially when both pathways are blocked; but most patients with most types of tumors still do not show a response. IDO is linked to both the CTLA-4 and PD-1 checkpoints via complex loops that are not yet fully elucidated. Thus, for example, CTLA-4 expression on Treg cells can up-regulate IDO in DCs [105], while IDO can reciprocally activate Treg cells. IDO also up-regulates PD-1 expression on Treg cells, which contributes to maintenance of PTEN activity [86]. Thus, IDO interacts with other important checkpoints. Mechanistically, however, both CTLA-4 and PD-1 are inhibitory checkpoints that are expressed on T cells; thus, they primarily come into play after antigen has been presented. IDO is somewhat different in that many of its actions occur further upstream, affecting the initial T cell activation step by altering the biology of the APC itself and the antigen-presenting milieu. Indeed, IDO appears to have certain inherent signaling functions in DCs that are distinct from its tryptophan-catabolizing enzymatic function [106], such that expression of IDO may render the DC more tolerogenic in a cell-intrinsic fashion [107]. Thus, due to its upstream effects on the antigen-presenting milieu, IDO may be non-redundant with the more distal T cell checkpoints, and blocking IDO concurrently with CTLA-4 or PD-1 might confer additive benefit. Preclinical models support this concept, both for CTLA-4 and for PD-1 [108–110], and clinical trials of these combinations are on-going (see below).
IDO and MDSCs

Myeloid-derived suppressor cells (MDSCs) are another important immunosuppressive cell population in the tumor microenvironment. To date, the relationship between IDO and MDSCs has not been well studied. In mouse models, MDSCs suppress T cells via production of reactive oxygen species, nitric oxide or arginase [111], but they do not appear to express IDO. However, IDO may contribute to recruitment of MDSCs into the tumor via more indirect mechanisms [41]; and local IDO and IDO-activated Tregs may increase the suppressor function of MDSCs once they reach the tumor [112]. In humans, several recent reports have described cells with a phenotype consistent with MDSCs, which also express high levels of immunosuppressive IDO [113-115]. Thus, IDO and MDSCs may have more interaction than previously thought, and further study is needed to define the relationship.

IDO and complement

Finally, the IDO pathway has recently been linked to control of the complement pathway in tumors [116]. The mechanism of this link is still not elucidated, but widespread complement activation was previously described when IDO was blocked during murine pregnancy [3]. In mouse brain tumors, the combination of radiation plus chemotherapy caused extensive complement deposition when IDO was blocked (but not when IDO was active), and this complement was mechanistically required for the effect of the IDO-inhibitor on survival [116]. Further studies are needed to elucidate the mechanism of this link between IDO and complement.

IDO-inhibitors in the clinic

Mechanistically, IDO-inhibitor drugs do not kill tumor cells directly, nor do they spontaneously initiate an immune response by themselves. Thus, the role for these drugs in the clinic will likely be to allow and enhance the immune responses triggered chemotherapy, vaccine or checkpoint inhibitors. Fortunately, in most studies the toxicity of IDO-inhibitor drugs appears low, even with sustained administration, so combinations should be feasible. Table 1 gives a partial listing of on-going clinical trials of IDO-inhibitor drugs, grouped by combination with chemotherapy or immune-modulator. Two agents currently account for the majority of the trials: indoximod (1-methyl-D-tryptophan), an inhibitor of the IDO pathway [85; 108-109; 112], and epacadostat (INCB024360) [110; 117-118]. No final results have yet been published, so it is too early to assess the benefits of these strategies, but anecdotal results have been encouraging. Additional IDO inhibitors are in the development pipeline, as well as agents that may target IDO2 [119] or TDO [120-121], two other tryptophan-catabolizing enzymes that may affect tumor immunity [122].

In monitoring clinical trials of IDO-inhibitors, target-validation studies and pharmacodynamic markers have been somewhat of a challenge. For target detection, there is no accepted set of cross-validated monoclonal antibodies for measuring IDO in clinical biopsy samples. This has made it hard to compare reports from different groups. The issue is further complicated by the fact that IDO is inducible, so the initial (pre-treatment) biopsy may not reflect the status during therapy. Pharmacodynamic markers have also been problematic. Kynurenine, the first metabolic product of IDO, is constitutively present in
plasma at low levels, because it is produced by TDO in the liver and homeostatic IDO in the gut and elsewhere [123]. In a subset of cancer patients, plasma kynurenine may be elevated by the presence of tumors (ref. [124] and references therein); and in these patients, if kynurenine returns to normal when treated with a with an IDO-inhibitor, then kynurenine is a useful marker of therapy. But in the majority of patients whose kynurenine is already in the normal range to start with, interpretation of kynurenine as a biomarker has been difficult. Target detection and biomarkers are areas that will need additional development as IDO-inhibitors enter clinical trials.

Concluding remarks

The field of cancer immunotherapy has recently entered a more mature phase. It has now become possible to achieve impressive, clinically-meaningful responses in at least a subset of selected patients. Now the question is how to enhance these responses, broaden them, and leverage the power of the patients’ own immune system to sustain them. IDO is potentially an attractive target for combinatorial immunotherapy because it affects inflammation, antigen cross-presentation, and the overall tolerogenic milieu. These are areas that are not directly targeted by current existing immunotherapy strategies, so there may be opportunities for synergy. However, more research is needed on a number of open issues (see Outstanding Questions).

The network of immunosuppressive mechanisms in the tumor microenvironment is complex, multifactorial and mutually reinforcing. It is based largely on a pathologic exaggeration of existing, natural regulatory processes in the immune system – especially mechanisms for the counter-regulatory control of inflammation, and for induction of acquired peripheral tolerance. Elucidating the role of IDO in these pathways can help to understand the abnormal biology of the tumor, and identify the most effective way to target IDO for therapy.

References


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**Trends Box**

- IDO can be induced in the tumor microenvironment by the spontaneous inflammation and T cell activation associated with many tumors; however, IDO may also be induced reactively, in response to inflammation induced by anti-tumor immunotherapy.

- IDO is rapidly induced when mice are challenged with apoptotic cells. Inhibiting or genetically deleting IDO results in loss of self-tolerance to apoptotic cell-associated antigens, and susceptibility to lupus-like autoimmunity.

- DNA-containing nanoparticles can induce immunosuppressive IDO via a pathway dependent on the STING sensor of cytosolic DNA.

- When Tregs cells are activated in the presence of IDO, they up-regulate a highly suppressive phenotype driven by the PTEN lipid phosphatase. In vivo, challenge with apoptotic cells triggers IDO-mediated induction of PTEN-expressing Tregs cells.

- In tumors, IDO induction by apoptotic cells may become relevant as the tumor attempts to inhibit immune responses to dying tumor cells following chemotherapy or immunotherapy.
**Outstanding Questions**

- How do tumors recruit immune checkpoints such as IDO during their development? Is this a natural response to the inflammatory signals associated with malignant transformation, dying cells and DNA release? Or are there in addition, tumor-induced processes that exaggerate the amount of IDO induced by this inflammation? And, in a related question, what is the relative contribution of IDO expressed in immune cells and stromal cells versus that expressed by tumor cells?

- What is the role of IDO in enforcing tolerance to apoptotic tumor cells? Is IDO induced in the tumor microenvironment (or tumor-draining lymph nodes) by dying tumor cells after therapy? Does this elicit PTEN-expressing Treg cells, and thereby suppress the response to immunogenic tumor antigens? And, if so, can IDO-inhibitor drugs be used to promote epitope-spreading and activation of new, endogenous T cells after chemotherapy or immunotherapy?

- What are the other relevant immunosuppressive mechanisms that must be blocked in order to maximize the effects of blocking IDO? Inhibiting IDO is not a magic bullet, and it seems to be a reproducible theme that IDO-inhibitors work better in combination with CTLA-4 or PD 1 blockade, or with some form of chemotherapy or adoptive-transfer. But what are the mechanisms, and what is the most rational basis for combinations with other agents?
Figure 1. The Tipping Point: is DNA sensed to impede or incite tumorigenesis?
A. During tumor growth DNA from dying cells is sensed by cGAS to stimulate STING/IFNαβ signaling, which potentiates immunogenic responses if tumors have relatively high antigenicity. These responses lead to tumor regression mediated by cytotoxic (CTL) and helper (Th) T cells. B. DNA is sensed to activate STING/IFNαβ signaling and induce IDO during tumorigenesis. IDO mediates dominant tolerogenic responses that activate Tregs to overcome immunogenic responses and promote tumorigenesis. These responses are more likely when tolerance thresholds are high due to low tumor antigenicity.
During initial TCR-mediated activation of resting Tregs, signals via the Akt and mTOR pathways can potentially destabilize the Tregs and cause reprogramming into a pro-inflammatory helper-like phenotype (“ex-Tregs”). To prevent this, some signal must inhibit the mTOR/Akt axis during activation. If the antigen-presenting cell expresses IDO (e.g., due to up-regulation by apoptotic cells or other inflammatory signals) then this can inhibit mTORC2 signaling via a process mediated by low tryptophan and GCN2 kinase. In principle, depletion of other amino acids such as arginine (by local Arginase I) could create a similar GCN2-mediated inhibition of mTORC2. Potentially, other metabolic stresses in the tumor microenvironment might likewise affect mTORC1 and the feedback loop to mTORC2 and Akt. By whatever pathway, if Akt phosphorylation on the activating Ser473 residue is blocked, then the Treg is able to maintain expression of FoxO3a and acquire a highly suppressive phenotype that includes up-regulation of cell-surface PD-1 and the lipid phosphatase PTEN. PD-1 can signal via PTEN to inhibit PI3K activity, and thus block phosphorylation of the activating Thr308 residue on Akt. This self-sustaining feedback loop acts to stably maintain the inhibition of Akt long-term, even if the original IDO or other metabolic stress is removed. Neuropilin-1 (Nrp-1) can also activate PTEN, and thus may be able to establish a similar stable activation state in the Treg.
Figure 3. Potential model of tolerogenic versus immunogenic cell death in the tumor microenvironment

Tumor cells undergo a wave of cell death in response to chemotherapy, radiation or immunotherapy (e.g., T cell adoptive transfer, or other immunotherapy that activates cytotoxic T cells, CTLs). Some of the tumor cells may die via an immunogenic pathway that releases pro-inflammatory mediators such as HMGB1, ATP and STING. This immunogenic cell death has the potential to trigger inflammation, immunogenic cross-presentation of tumor antigens, and robust T cell activation. However, some fraction of the tumor cells also likely die by classical apoptosis, with consequent induction of immunosuppressive pathways such as TGFβ, IDO and IDO-induced PTEN-Treg activation. Unless the pro-inflammatory signals are very strong, these immunosuppressive signals (particularly activation of the potent PTEN-Tregs) are likely to be dominant. Thus, the net outcome of tumor cell death is often immune suppression and tolerance. Underneath, however, the immunogenic cell death may still be present, and available to drive anti-tumor immune responses if the dominant suppression is blocked.
### Table 1

Clinical trials of IDO-inhibitors

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<th>Strategy</th>
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